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(54) Title: CHIMERIC ADENOVIRAL VECTORS

(57) Abstract

A chimeric adenoviral vector is provided that comprises nucleotide sequence of a first adenovirus, wherein all or part of at least one gene of said first adenovirus encoding a protein that facilitates binding of said vector to a target mammalian cell, or internalization thereof within said cell, is replaced by all or part of the corresponding gene from a second adenovirus belonging to subgroup D, said vector further comprising a transgene operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell. Compositions comprising such vectors and methods of using such vectors to deliver transgenes to target mammalian cells, particularly airway epithelial cells, are also provided.

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Description

Chimeric Adenoviral Vectors

5 Introduction

The present invention relates to chimeric adenoviral vectors, that is, vectors comprising DNA from more than one scrotype of adenovirus, which offer enhanced infection efficiency of target cells in order to deliver one or more therapeutically useful nucleotide sequences, including transgenes, therein. Such a nucleotide sequence may comprise a gene not otherwise present in the target cell that codes for a therapeutic and/or biologically active protein, or may represent, for example, an active copy of a gene that is already present in the target cell, but in a defective or deficient form.

15 Background of the Invention

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One of the fundamental challenges now facing medical practicioners is that although the defective genes that are associated with numerous inherited diseases (or that represent disease risk factors including for various cancers) have been isolated and characterized, methods to correct the disease states themselves by providing patients with normal copies of such genes (the technique of gene therapy) are substantially lacking. Accordingly, the development of improved methods of intracellular delivery therefor is of great medical importance. Examples of diseases that it is hoped can be treated by gene therapy include inherited disorders such as cystic fibrosis, Gaucher's disease, Fabry's disease, and muscular dystrophy.

Representative of acquired disorders that can be treated are: (1) for cancers: multiple myeloma, leukemias, melanomas, ovarian carcinoma and small cell lung cancer; (2) for cardiovascular conditions: progressive heart failure, restenosis, and hemophilias; and (3) for neurological conditions: traumatic brain injury.

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Gene therapy requires successful transfer of nucleic acid to the target cells of a patient. Gene transfer may generally be defined as the process of introducing an expressible polynucleotide (for example a gene, a cDNA, or an mRNA patterned thereon) into a cell. In a particular application of this approach, successful expression of an encoding polynucleotide leads to production in the cells of a normal protein and leads to correction of a disease state associated with an abnormal gene. Therapies based on providing such proteins directly to target cells (protein replacement therapy) have generally proved ineffective since, for example, the cell membrane presents a selectively permeable barrier to entry. Thus there is great interest in alternative methods to cause delivery of therapeutic proteins, especially by transfer of the relevant polynucleotide, often referred to as a transgene.

Viral vectors have been used with increasing frequency to date to deliver transgenes to target cells. Most attempts to use viral vectors for gene therapy have relied on retrovirus-based vectors, chiefly because of their ability to integrate into the cellular genome. However, the disadvantages of retroviral vectors are becoming increasingly clear, including their tropism for dividing cells only, the possibility of insertional mutagenesis upon integration into the cell genome, decreased expression of the transgene over time, rapid inactivation by serum complement, and the possibility of generation of replication-competent retroviruses. See, for example, D. Jolly, et al., Cancer Gene Therapy, 1, 1994, pp. 51-64, and C.P. Hodgson, et al., Bio Technology, 13, 1995, pp. 222-225. Such disadvantages have led to the development of other viral-based vector systems, including those derived from adenoviruses.

Adenovirus (Ad) is a nuclear DNA virus with a genome of about 36 kb, which has been well-characterized through studies in classical genetics and molecular biology. A detailed discussion of adenovirus is found in Thomas Shenk, "Adenoviridae and their Replication", and M. S. Horwitz, "Adenoviruses", Chapters 67 and 68, respectively, in Virology, B.N. Fields et al., eds., 2nd edition, Raven Press, Ltd., New York, 1996, and reference therein is found to numerous aspects of adenovirus pathology, epidemiology, structure, replication, genetics and classification.

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In a simplified form, the adenoviral genome is classified into early (known as E1-E4) and late (known as L1-L5) transcriptional units, referring to the generation of two temporal classes of viral proteins. The demarcation between these events is viral DNA replication.

The human adenoviruses are divided into numerous scrotypes (approximately 47, numbered accordingly and classified into 6 subgroups: A, B, C, D, E and F), based upon properties including hemagglutination of red blood cells, oncogenicity, DNA base and protein amino acid compositions and homologies, and antigenic relationships. Additional background information concerning Ad serotype classification, including that for subgroup D, can be found, for example, in F. Deryckere et al., Journal of Virology, 70, 1996, pp. 2832-2841; and A. Bailey et al., Virology, 205, 1994, pp. 438-452, and in other art-recognized references.

Adenoviruses are nonenveloped, regular icosahedrons (having 20 triangular surfaces and 12 vertices) that are about 65-80 nm in diameter. A protein called fiber projects from each of these vertices. The fiber protein is itself generally composed of 3 identical polypeptide chains, although the length thereof varies between serotypes. The protein coat (capsid) is composed of 252 subunits (capsomeres), of which 240 are hexons, and 12 are pentons. Each penton comprises a penton base, on the surface of the capsid, and a fiber protein projecting from the base. The Ad 2 penton base protein, for example, has been determined to be a 8 x 9 nm ring shaped complex composed of 5 identical protein subunits of 571 amino acids each.

Current understanding of adenovirus-cell interactions suggests that adenovirus utilizes two cellular receptors to attach to, and then infect a target cell. It has been further suggested that the fiber protein of an infecting adenovirus first attaches to a receptor, the identity of which is still unknown, and then penton base attaches to a further receptor, often a protein of the alpha integrin family. It has been determined that alpha-integrins often recognize short amino acid sequences on other cellular proteins for attachment pruposes including the tripeptide sequence Arg-Gly-Asp (abbreviated RGD). An RGD sequence is also found in the penton base protein of

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adenovirus and is currently understood in the art to mediate attachment of Ad to alpha integrins.

Recombinant adenoviruses have several advantages for use as gene transfer vectors, including tropism for both dividing and non-dividing cells, minimal pathogenic potential, ability to replicate to high titer for preparation of vector stocks, and the potential to carry large inserts (Berkner, K.L., Curr. Top. Micro. Immunol. 158:39-66, 1992; Jolly, D., Cancer Gene Therapy 1:51-64, 1994).

The carrying capacity of an adenovirus vector is proportional to the size of the adenovirus genome present in the vector. For example, a capacity of about 8 kb can be created from the deletion of certain regions of the virus genome dispensable for virus growth, e.g., E3, and the deletion of a genomic region such as E1 whose function may be restored in trans from 293 cells (Graham, F.L., J. Gen. Virol. 36:59-72, 1977) or A549 cells (Imler et al., Gene Therapy 3:75-84, 1996). Such E1-deleted vectors are rendered replication-defective, which is desirable for the engineering of adenoviruses for gene transfer. The upper limit of vector DNA capacity for optimal carrying capacity is about 105%-108% of the length of the wild-type genome. Further adenovirus genomic modifications are possible in vector design using cell lines which supply other viral gene products in trans, e.g., complementation of E2a (Zhou et al., J. Virol. 70:7030-7038, 1996), complementation of E4 (Krougliak et al., Hum. Gene Ther. 6:1575-1586, 1995; Wang et al., Gene Ther. 2:775-783, 1995), or complementation of protein IX (Caravokyri et al., J. Virol. 69:6627-6633, 1995; Krougliak et al., Hum. Gene Ther. 6:1575-1586, 1995). Maximal carrying capacity can be achieved using adenoviral vectors deleted for all viral coding sequences (Kochanek et al., Proc. Natl. Acad. Sci. USA 93:5731-5736, 1996; Fisher et al., Virology 217:11-22, 1996).

Transgenes that have been expressed to date by adenoviral vectors include p53 (Wills et al., Human Gene Therapy 5:1079-188, 1994); dystrophin (Vincent et al., Nature Genetics 5:130-134, 1993; erythropoietin (Descamps et al., Human Gene Therapy 5:979-985, 1994; ornithine transcarbamylase (Stratford-Perricaudet et al.,

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Human Gene Therapy 1:241-256, 1990; We et al., J. Biol. Chem. 271;3639-3646, 1996;); adenosine deaminase (Mitani et al., Human Gene Therapy 5:941-948, 1994); interleukin-2 (Haddada et al., Human Gene Therapy 4:703-711, 1993); and α1-antitrypsin (Jaffe et al., Nature Genetics 1:372-378, 1992); thrombopoietin (Ohwada et al., Blood 88:778-784, 1996); and cytosine deaminase (Ohwada et al., Hum. Gene Ther. 7:1567-1576, 1996).

The particular tropism of adenoviruses for cells of the respiratory tract has particular relevance to the use of adenovirus in gene therapy for cystic fibrosis (CF), which is the most common autosomal recessive disease in Caucasians. The disease is caused by the presence of one or more mutations in the gene that encodes a protein known as cystic fibrosis transmembrane conductance regulator (CFTR), and which regulates the movement of ions (and therefore fluid) across the cell membrane of epithelial cells, including lung epithelial cells. Abnormal ion transport in airway cells leads to abnormal mucous secretion, inflammmation and infection, tisssue damage, and eventually death. Mutations in the CFTR gene that disturb the cAMP-regulated Cl channel in airway epithelia result in pulmonary dysfunction (Zabner et al., Nature Genetics 6:75-83, 1994). Adenovirus vectors engineered to carry the CFTR gene have been developed (Rich et al., Human Gene Therapy 4:461-476, 1993) and studies have shown the ability of these vectors to deliver CFTR to nasal epithelia of CF patients (Zabner et al., Cell 75:207-216, 1993), the airway epithelia of cotton rats and primates (Zabner et al., Nature Genetics 6:75-83, 1994), and the respiratory epithelium of CF patients (Crystal et al., Nature Genetics 8:42-51, 1994). Recent studies have shown that administering an adenoviral vector containing a DNA sequence encoding CFTR to airway epithelial cells of CF patients can restore a functioning chloride ion channel in the treated epithelial cells (Zabner et al., J. Clin. Invest. 97:1504-1511, 1996; U.S. Patent No. 5,670,488 issued September 23, 1997).

Serotype classification is partly based on viral surface protein sequence variation. Because the infectious capabilities of the virus are associated with the surface protein interactions of the virus with cellular proteins, the serotype is an

important determinant of viral entry into target cells, and can account for the infectious heterogeneity of adenovirus serotypes. Most adenoviral vectors have been constructed using adenovirus serotypes from the well-studied group C adenoviruses, especially Ad 2 and Ad 5. However, other adenovirus serotypes display infectious properties that are relevant to the further design of improved adenoviral vectors, for example, those derived from subgroup D, which display enhanced tropism for human airway epithelial cells.

It is widely hoped that gene therapy will provide a long lasting and predictable form of therapy for certain disease states, and it is likely the only form of therapy suitable for many inherited diseases. Although adenoviral vectors are currently in clinical use and have shown therapeutic promise, a need remains to improve the infection efficiency of these vectors in order to further improve their gene transfer capabilities. The present invention addresses this goal.

15 Summary Of The Invention

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The present invention provides for chimeric adenoviral vectors which offer enhanced infection efficiency of target cells for the delivery of one or more transgenes. In a representative aspect of the invention, the vectors comprise nucleotide sequences coding for therapeutically useful proteins and have enhanced tropism for airway epithelial cells.

Accordingly, there are provided chimeric adenoviral vectors comprising nucleotide sequence of a first adenovirus, wherein at least one gene of said first adenovirus encoding a protein that facilitates binding of said vector to a target mammalian cell, or internalization thereof within said cell, is replaced by the corresponding gene from a second adenovirus belonging to subgroup D. These vectors may further comprising a transgene operably linked to a eucaryotic promoter or other regulatory elements to allow for expression therefrom in a mammalian cell. In a representative aspect thereof, the replaced encoding sequence codes for Ad fiber, hexon or penton base.

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In a further preferred embodiment of the invention, there are provided chimeric adenoviral vectors comprising nucleotide sequence of a first adenovirus, wherein a portion of a gene thereof encoding a protein that facilitates binding of said vector to a target mammalian cell, or internalization thereof within said cell, is replaced by a portion of the corresponding gene from a second adenovirus belonging to subgroup D. These vectors may further comprising a transgene operably linked to a eucaryotic promoter or other regulatory elements to allow for expression therefrom in a mammalian cell. In a representative aspect thereof, the replaced encoding sequence codes for a portion of Ad fiber, hexon or penton base.

Preferably, the second adenovirus is a member of subgroup D, and the replaced nucleotide sequence encodes a polypeptide selected from the group consisting of Ad fiber, a fragment of Ad fiber, Ad hexon, a fragment of Ad hexon, Ad penton base, and a fragment of Ad penton base. In a preferred embodiment, said second adenovirus is selected from the group consisting of serotypes Ad 9, Ad 15, Ad 17, Ad 19, Ad 20, Ad 22, Ad 26, Ad 27, Ad 28, Ad 30, and Ad 39. In preferred embodiments of the chimeric adenoviral vectors, the first adenovirus is selected from the group consisting of Ad 2, Ad 5, and Ad 12.

The invention is also directed to compositions comprising the chimeric adenoviral vectors of the invention. Additional aspects of the invention include methods to use the chimeric adenoviral vectors of the invention to deliver transgenes to mammalian target cells, for example, to the airway epithelial cells of patients.

A still further representative apsect of the invention involves a method of providing a therapeutic and/or biologically active protein to the airway epithelial cells of a patient by administering to said cells an adenoviral vector comprising elements of an Ad 17 genome, and a transgene encoding said therapeutic protein that is operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell, under conditions whereby the transgene encoding said therapeutic protein is expressed, and therapeutic benefit is produced in said airway epithelial cells.

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These and other aspects of the present invention are described in the Detailed Description of the Invention which follows directly.

Brief Description of the Drawings

FIGURE 1 depicts infection of NHBE cells by Ad 2.

FIGURE 2 depicts infection of NHBE cells by Ad 17.

FIGURE 3 plots the result of binding to human nasal polyp epithelial cell isolates by Ad 2 and Ad 17.

FIGURE 4 is a map of the vector Ad2/βgal-2/fiber Ad 17.

FIGURE 5 shows a comparison of the amino acid sequence of penton base from Ad 17 (top) [SEQ ID NO: 4] and Ad 2 (bottom) [SEQ ID NO: 5], and further depicts the variable RGD containing region.

FIGURE 6 depicts an amino acid sequence pileup for penton base from particular Ad serotypes, including f10 (from fowl) [SEQ ID NO: 6 through SEQ ID NO: 10].

FIGURE 7 shows a comparison of the amino acid sequence of fiber from Ad 17 (top) [SEQ ID NO: 11] and Ad 2 (bottom) [SEQ ID NO: 12].

FIGURE 8 depicts an amino acid sequence pileup for fiber from particular Ad serotypes [SEQ ID NO: 11 through SEQ ID NO: 22], including two forms of serotype 40 (40-1 and 40-2) which differ in that one variant has two (but non-identical) copies of the fiber gene.

FIGURE 9 shows the infection efficiency of colon cancer cell lines by adenovirus serotypes.

FIGURE 10 shows the infection efficiency of cancer cell lines by adenovirus serotypes.

Provided in the Sequence Listing attached hereto are also:

SEQ ID NO: 1, the complete nucleotide sequence of Ad 17;

SEQ ID NO: 2, the complete encoding nucleotide sequence for Ad 17 fiber;

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SEQ ID NO: 3, the complete encoding nucleotide sequence for Ad 17 penton base.

Detailed Description of the Invention

The present invention provides for chimeric adenoviral vectors comprising nucleotide sequence of a first adenovirus, wherein at least one gene of said first adenovirus encoding a protein that facilitates binding of said vector to a target mammalian cell, or internalization thereof within said cell, is replaced by the corresponding gene from a second adenovirus belonging to subgroup D, said vectors further comprising a transgene operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell. In a representative aspect thereof, the replaced encoding sequence correspond to the gene encoding the Ad fiber, hexon or penton base proteins, or combinations thereof.

In a further preferred embodiment of the invention, there are provided chimeric adenoviral vectors comprising nucleotide sequence of a first adenovirus, wherein a portion of a gene thereof encoding a protein that facilitates binding of said vector to a target mammalian cell, or internalization thereof within said cell, is replaced by a portion of the corresponding gene from a second adenovirus belonging to subgroup D, said vectors further comprising a transgene operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell. In a representative aspect thereof, the replaced encoding sequence codes for a portion of the Ad fiber, hexon or penton base proteins, or combinations thereof. Where a portion of a gene from a second adenovirus is used to construct a chimeric adenoviral vector, such sequence will have a length sufficient to confer a desired serotypic-specific virus-cell interaction to the vector.

The present invention involves the recognition that adenoviral vectors that are either based substantially upon the genome of Ad serotypes classified in subgroup D, or that contain certain Ad-protein encoding polynucleotide sequences of subgroup D adenovirus, are particularly effective at binding to, and internalizing within, human

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cells, such that therapeutic transgenes included in the adenoviral vector are efficiently expressed. This discovery is particularly surprising given that adenovirus serotypes of subgroup D are not clinically associated with human respiratory disease, and that, for example association with conjunctivitis is more typical. The recognition of this tropism is of particular relevance for the treatment by gene therapy of recognized disease states such as cystic fibrosis or α 1-antitrypsin deficiency. This discovery is particularly surprising given that adenovirus serotypes of subgroup D are not clinically associated with human respiratory disease, and that, for example association with conjunctivitis is more typical. The recognition of this tropism is of particular relevance for the treatment by gene therapy of recognized disease states such as cystic fibrosis or α 1-antitrypsin deficiency.

In a representative aspect of the invention, the adenoviral vectors further comprise nucleotide sequences coding for one or more transgenes and have enhanced tropism for airway epithelial cells. Preferably, the chimeric adenoviral vectors are replication-defective, a feature which contributes to the enhanced safety of adenoviral vectors administered to individuals.

Preferably, the second adenovirus is a member of subgroup D, and the replaced nucleotide sequence encodes a polypeptide selected from the group consisting of Ad fiber, a fragment of Ad fiber, Ad hexon, a fragment of Ad hexon, Ad penton base, and a fragment of Ad penton base. In a preferred embodiment, said second adenovirus is selected from the group consisting of serotypes Ad 9, Ad 15, Ad 17, Ad 19, Ad 20, Ad 22, Ad 26, Ad 27, Ad 28, Ad 30, and Ad 39. In a most preferred embodiment, the second adenovirus is Ad 17. In other preferred embodiments of the chimeric adenoviral vectors, the first adenovirus is selected from the group consisting of Ad 2, Ad 5, and Ad 12.

There is substantial evidence that any reported transforming properties of the E4 region of certain subgroup D serotypes do not extend to Ad serotypes whose use is preferred according to the practice of the present invention (see, for example, R. Javier

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et al., Science, 257, 1992, pp. 1267-1271). It is expected also that, for example, individual ORFs of subgroup D E4 region, such as ORF1, could be deleted.

Additional aspects of the invention include methods to provide biologically active and/or therapeutic proteins to mammalian cells, including, but not limited to, the airway epithelial cells of individuals, in order to provide phenotypic benefit. According to this aspect of the invention, chimeric adenoviral vectors are used in which a nucleotide sequence of a first adenovirus is replaced by the corresponding nucleotide sequence of a second adenovirus. Preferably, the second adenovirus is a member of subgroup D, and the replaced nucleotide sequence encodes a polypeptide encoding all or part of Ad fiber, Ad hexon, or Ad penton base, or combinations thereof.

A still further representative aspect of the invention involves providing a biologically active and/or therapeutic protein in the airway epithelial cells of a patient by administering to said cells an adenoviral vector comprising elements of an Ad 17 genome, and a transgene encoding said protein that is operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell, under conditions whereby the transgene encoding said protein is expressed, and the desired phenotypic benefit is produced in said airway epithelial cells. According to the practice of the invention, it is preferred that an chimeric adenovirus vector utilized to deliver a transgene to the respiratory epithelium (including that of the nasal airway, trachea, and bronchi and alveoli of the lung), or to other tissues of the body, comprise serotypes within subgroup D, as such classification is recognized in the art.

In order to construct the chimeric adenoviral vectors of the invention, reference may be made to the substantial body of literature on how such vectors may be designed, constructed and propagated using techniques from molecular biology and microbiology that are well-known to the skilled artisan. Specific examples of adenoviral vector genomes which can be used as the backbone for a chimeric adenoviral vector of the invention include, for example, Ad2/CFTR-1 and Ad2/CFTR-2 and others described in U. S. Patent No. 5,670,488, issued September 23, 1997

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(incorporated herein by reference). Such vectors may include deletion of the E1 region, partial or complete deletion of the E4 region, and deletions within, for example, the E2 and E3 regions. Within the scope of the invention are, for example, chimeric vectors which contain an Ad 2 backbone with one or more Ad 17 capsid proteins or fragments thereof in the virus. Other adenoviral vector genomic designs which can be used in the chimeric adenoviral vectors of the invention include those derived from allowed U.S. Patent Application Serial No. 08/409,874, filed March 24, 1995, and allowed U.S. Patent Application Serial No. 08/540,077, filed October 6, 1995 (both incorporated herein by reference).

To construct the recombinant chimeric adenoviral vectors of the invention which contain a transcription unit, the skilled artisan can use the standard techniques of molecular biology to engineer a transgene or a capsid protein into a backbone vector genome (Berkner, K.L., Curr. Top. Micro. Immunol. 158:39-66, 1992). For example, a plasmid containing a transgene and any operably linked regulatory elements inserted into an adenovirus genomic fragment can be co-transfected with a linearized viral genome derived from an adenoviral vector of interest into a recipient cell under conditions whereby homologous recombination occurs between the genomic fragment and the virus. Preferably, a transgene is engineered into the site of an E1 deletion. As a result, the transgene is inserted into the adenoviral genome at the site in which it was cloned into the plasmid, creating a recombinant adenoviral vector. The chimeric adenoviral vectors can also be constructed using standard ligation techniques, for example, removing a restriction fragment containing a fiber gene from a first adenovirus and ligating into that site a restriction fragment containing a fiber gene from a second adenovirus. A representative example of a chimeric adenoviral vector of the invention is Ad2/βgal-2 fiber 17 (exemplified in Example 6).

Construction of the chimeric adenoviral vectors can be based on adenovirus DNA sequence information widely available in the field, e.g., nucleic acid sequence databases such as GenBank.

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Preparation of replication-defective chimeric adenoviral vector stocks can be accomplished using cell lines that complement viral genes deleted from the vector, e.g., 293 or A549 cells containing the deleted adenovirus E1 genomic sequences. The use of HER3 cells (human embryonic retinoblasts transformed by Ad 12), as a complementing cell line is of note. After amplification of plaques in suitable complementing cell lines, the viruses can be recovered by freeze-thawing and subsequently purified using cesium chloride centrifugation. Alternatively, virus purification can be performed using chromatographic techniques, e.g., as set forth in International Application No. PCT/US96/13872, filed August 30, 1996, incorporated herein by reference.

Titers of replication-defective chimeric adenoviral vector stocks can be determined by plaque formation in a complementing cell line, e.g., 293 cells. Endpoint dilution using an antibody to the adenoviral hexon protein may be used to quantitate virus production or infection efficiency of target cells (Armentano et al., Hum. Gene Ther. 6:1343-1353, 1995, incorporated herein by reference).

Transgenes which can be delivered and expressed from a chimeric adenoviral vector of the invention include, but are not limited to, those encoding enzymes, blood derivatives, hormones, lymphokines such as the interleukins and interferons, coagulants, growth factors, neurotransmitters, tumor suppressors, apoliproteins, antigens, and antibodies, and other biologically active proteins. Specific transgenes which may be encoded by the chimeric adenoviral vectors of the invention include, but are not limited to, cystic fibrosis transmembrane regulator (CFTR), dystrophin, glucocerebrosidase, tumor necrosis factor, p53, p21, herpes simplex thymidine kinase and gancyclovir, retinoblastoma (Rb), and adenosine deaminase (ADA). Transgenes encoding antisense molecules or ribozymes are also within the scope of the invention. The vectors may contain one or more transgenes under the control of one or more regulatory elements.

In addition to containing the DNA sequences encoding one or more transgenes, the chimeric adenoviral vectors of the invention may contain any

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expression control sequences such as a promoter or enhancer, a polyadenylation element, and any other regulatory elements that may be used to modulate or increase expression, all of which are operably linked in order to allow expression of the transgene. The use of any expression control sequences, or regulatory elements, which facilitate expression of the transgene is within the scope of the invention. Such sequences or elements may be capable of generating tissue-specific expression or be susceptible to induction by exogenous agents or stimuli.

Infection of target cell by the chimeric adenoviral vectors of the invention may also be facilitated by the use of cationic molecules, such as cationic lipids as disclosed in PCT Publication No. WO96/18372, published June 20, 1996, incorporated herein by reference.

Cationic amphiphiles have a chemical structure which encompasses both polar and non-polar domains so that the molecule can simultaneously facilitate entry across a lipid membrane with its non-polar domain while its cationic polar domain attaches to a biologically useful molecule to be transported across the membrane.

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Cationic amphiphiles which may be used to form complexes with the chimeric adenoviral vectors of the invention include, but are not limited to, cationic lipids, such as DOTMA (Felgner et al., Proc. Natl. Acad. Sci. USA 84:7413-7417, 1987) (N-[1-(2,3-dioletloxy)propyl]-N,N,N - trimethylammonium chloride); DOGS 20 (dioctadecylamidoglycylspermine) (Behr et al., Proc. Natl. Acad. Sci. USA 86:6982-6986, 1989); DMRIE (1,2-dimyristyloxypropyl-3-dimethyl-hydroxyethyl ammonium bromide) (Felgner et al., J. Biol. Chem. 269:2550-2561, 1994; and DC-chol (3B [N-N', N'-dimethylaminoethane) -carbamoyl] cholesterol) (U.S. Patent No. 5, 283,185 to Epand et al.). The use of other cationic amphiphiles recognized in the art or which come to be discovered is within the scope of the invention.

In preferred embodiments of the invention, the cationic amphiphiles useful to complex with and facilitate transfer of the vectors of the invention are those lipids which are described in PCT Publication No. WO96/18372, published June 20, 1996, which is incorporated herein by reference. Preferred cationic amphiphiles described

herein to be used in the delivery of the plasmids and/or viruses are GL-53, GL-67, GL-75, GL-87, GL-89, and GL-120, including protonated, partially protonated, and deprotonated forms thereof. Further embodiments include the use of non-T-shaped amphiphiles as described on pp. 22-23 of the aforementioned PCT application, including protonated, partially protonated and deprotonated forms thereof. Most preferably, the cationic amphiphile which can be used to deliver the vectors of the invention is spermine cholesterol carbamate (GL-67).

In the formulation of compositions comprising the chimeric adenoviral vectors of the invention, one or more cationic amphiphiles may be formulated with neutral colipids such as dileoylphosphatidylethanolamine (DOPE) to facilitate delivery of the vectors into a cell. Other co-lipids which may be used in these complexes include, but are not limited to, diphytanoylphosphatidylethanolamine, lysophosphatidylethanolamines, other phosphatidylethanolamines, phosphatidylcholines, lyso-phosphatidylcholines and cholesterol. A preferred molar ratio of cationic amphiphile to colipid is 1:1. However, it is within the scope of the invention to vary this ratio, including also over a considerable range. In a preferred embodiment of the invention, the cationic amphiphile GL-67 and the neutral co-lipid DOPE are combined in a 1:2 molar ratio, respectively, before complexing with a chimeric adenoviral vector for delivery to a cell.

In the formulation of complexes containing a cationic amphiphile with a chimeric adenoviral vector, a preferred range of 10^7 - 10^{10} infectious units of virus may be combined with a range of 10^4 - 10^6 cationic amphiphile molecules/viral particle.

The infection efficiency of the chimeric adenoviral vectors of the invention

may be assayed by standard techniques to determine the infection of target cells. Such methods include, but are not limited to, plaque formation, end-point dilution using, for example, an antibody to the adenoviral hexon protein, and cell binding assays using radiolabelled virus. Improved infection efficiency may be characterized as an increase in infection of at least an order of magnitude with reference to a control virus. Where

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a chimeric adenoviral vector encodes a marker or other transgene, relevant molecular assays to determine expression include the measurement of transgene mRNA, by, for example, Northern blot, S1 analysis or reverse transcription-polymerase chain reaction (RT-PCR). The presence of a protein encoded by a transgene may be detected by Western blot, immunoprecipitation, immunocytochemistry, or other techniques known to those skilled in the art. Marker-specific assays can also be used, such as X-gal staining of cells infected with a chimeric adenoviral vector encoding β -galactosidase.

In order to determine transgene expression and infection efficiency in vivo using the constructs and compositions of the invention, animal models may be particularly relevant in order to assess transgene persistence against a background of potential host immune response. Such a model may be chosen with reference to such parameters as ease of delivery, identity of transgene, relevant molecular assays, and assessment of clinical status. Where the transgene encodes a protein whose lack is associated with a particular disease state, an animal model which is representative of the disease state may optimally be used in order to assess a specific phenotypic result and clinical improvement. However, it is also possible that particular chimeric adenoviral vectors of the invention display enhanced infection efficiency only in human model systems, e.g., using primary cell cultures, tissue explants, or permanent cell lines. In such circumstances where there is no animal model system available in which to model the infection efficiency of a chimeric adenoviral vector with respect to human cells, reference to art-recognized human cell culture models will be most relevant and definitive.

Relevant animals in which the chimeric adenoviral vectors may be assayed include, but are not limited to, mice, rats, monkeys, and rabbits. Suitable mouse strains in which the vectors may be tested include, but are not limited to, C3H, C57Bl/6 (wild-type and nude) and Balb/c (available from Taconic Farms, Germantown, New York).

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Where it is desirable to assess the host immune response to vector administration, testing in immune-competent and immune-deficient animals may be

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compared in order to define specific adverse responses generated by the immune system. The use of immune-deficient animals, e.g., nude mice, may be used to characterize vector performance and persistence of transgene expression, independent of an acquired host response.

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In a particular embodiment where the transgene is the gene encoding cystic fibrosis transmembrane regulator protein (CFTR) which is administered to the respiratory epithelium of test animals, expression of CFTR may be assayed in the lungs of relevant animal models, for example, C57Bl/6 or Balb/c mice, cotton rats, or Rhesus monkeys. Molecular markers which may used to determine expression include the measurement of CFTR mRNA, by, for example, Northern blot, S1 analysis or RT-PCR. The presence of the CFTR protein may be detected by Western blot, immunoprecipitation, immunocytochemistry, or other techniques known to those skilled in the art. Such assays may also be used in tissue culture where cells deficient in a functional CFTR protein and into which the chimeric adenoviral vectors have been introduced may be assessed to determine the presence of functional chloride ion channels - indicative of the presence of a functional CFTR molecule.

The chimeric adenoviral vectors of the invention have a number of in vivo and in vitro utilities. The vectors can be used to transfer a normal copy of a transgene encoding a biologically active protein to target cells in order to remedy a deficient or dysfunctional protein. The vectors can be used to transfer marked transgenes (e.g., containing nucleotide alterations) which allow for distinguishing expression levels of a transduced gene from the levels of an endogenous gene. The chimeric adenoviral vectors can also be used to define the mechanism of specific viral protein-cellular protein interactions that are mediated by specific virus surface protein sequences. The vectors can also be used to optimize infection efficiency of specific target cells by adenoviral vectors, for example, using a chimeric adenoviral vector containing Ad 17 fiber protein to infect human nasal polyp cells. Where it is desirable to use an adenoviral vector for gene transfer to cancer cells in an individual, a chimeric adenoviral vector can be chosen which selectively infects the specific type of target

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cancer cell and avoids promiscuous infection. Where primary cells are isolated from a tumor in an individual requiring gene transfer, the cells may be tested against a panel of chimeric adenoviral vectors to select a vector with optimal infection efficiency for gene delivery. The vectors can further be used to transfer tumor antigens to dendritic cells which can then be delivered to an individual to elicit an anti-tumor immune response. Chimeric adenoviral vectors can also be used to evade undesirable immune responses to particular adenovirus serotypes which compromise the gene transfer capability of adenoviral vectors.

The present invention is further directed to compositions containing the chimeric adenoviral vectors of the invention which can be administered in an amount effective to deliver one or more desired transgenes to the cells of an individual in need of such molecules and cause expression of a transgene encoding a biologically active protein to achieve a specific phenotypic result. The cationic amphiphile-plasmid complexes or cationic amphiphile-virus complexes may be formulated into compositions for administration to an individual in need of the delivery of the transgenes.

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The compositions can include physiologically acceptable carriers, including any relevant solvents. As used herein, "physiologically acceptable carrier" includes any and all solvents, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents, and the like. Except insofar as any conventional media or agent is incompatible with the active ingredient, its use in the compositions is contemplated.

Routes of administration for the compositions containing the chimeric adenoviral vectors of the invention include conventional and physiologically acceptable routes such as direct delivery to a target organ or tissue, intranasal, intravenous, intramuscular, subcutaneous, intradermal, oral and other parenteral routes of administration.

The invention is further directed to methods for using the compositions of the invention in vivo or ex vivo applications in which it is desirable to deliver one or more

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transgenes into cells such that the transgene produces a biologically active protein for a normal biological or phenotypic effect. In vivo applications involve the direct administration of one ore more chimeric adenoviral vectors formulated into a composition to the cells of an individual. Ex vivo applications involve the transfer of a composition containing the chimeric adenoviral vectors directly to autologous cells which are maintained in vitro, followed by readministration of the transduced cells to a recipient.

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Dosage of the chimeric adenoviral vector to be administered to an individual for expression of a transgene encoding a biologically active protein and to achieve a specific phenotypic result is determined with reference to various parameters, including the condition to be treated, the age, weight and clinical status of the individual, and the particular molecular defect requiring the provision of a biologically active protein. The dosage is preferably chosen so that administration causes a specific phenotypic result, as measured by molecular assays or clinical markers. For example, determination of the infection efficiency of a chimeric adenoviral vector containing the CFTR transgene which is administered to an individual can be performed by molecular assays including the measurement of CFTR mRNA, by, for example, Northern blot, S1 or RT-PCR analysis or the measurement of the CFTR protein as detected by Western blot, immunoprecipitation, immunocytochemistry, or other techniques known to those skilled in the art. Relevant clinical studies which could be used to assess phenotypic results from delivery of the CFTR transgene include PFT assessment of lung function and radiological evaluation of the lung. Demonstration of the delivery of a transgene encoding CFTR can also be demonstrated by detecting the presence of a functional chloride channel in cells of an individual with cystic fibrosis to whom the vector containing the transgene has been administered (Zabner et al., J. Clin. Invest. 97:1504-1511, 1996). Transgene expression in other disease states can be assayed analogously, using the specific clinical parameters most relevant to the condition.

Dosages of a chimeric adenoviral vector which are effective to provide expression of a transgene encoding a biologically active protein and achieve a specific phenotypic result range from approximately 10⁸ infectious units (I.U.) to 10¹¹ I.U. for humans.

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It is especially advantageous to formulate parenteral compositions in dosage unit form for ease of administration and uniformity of dosage. Dosage unit form as used herein refers to physically discrete units suited as unitary dosages for the subjects to be treated, each unit containing a predetermined quantity of active ingredient calculated to produce the specific phenotypic effect in association with the required physiologically acceptable carrier. The specification for the novel dosage unit forms of the invention are dictated by and directly depend on the unique characteristics of the chimeric adenoviral vector and the limitations inherent in the art of compounding. The principal active ingredient (the chimeric adenoviral vector) is compounded for convenient and effective administration in effective amounts with the physiologically acceptable carrier in dosage unit form as discussed above.

Maximum benefit and achievement of a specific phenotypic result from administration of the chimeric adenoviral vectors of the invention may require repeated administration. Such repeated administration may involve the use of the same chimeric adenoviral vector, or, alternatively, may involve the use of different chimeric adenoviral vectors which are rotated in order to alter viral antigen expression and decrease host immune response.

The practice of the invention employs, unless otherwise indicated, conventional techniques of protein chemistry, molecular virology, microbiology, recombinant DNA technology, and pharmacology, which are within the skill of the art. Such techniques are explained fully in the literature. See, e.g., Current Protocols in Molecular Biology, Ausubel et al., eds., John Wiley & Sons, Inc., New York, 1995, and Remington's Pharmaceutical Sciences, 17th ed., Mack Publishing Co., Easton, PA, 1985.

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The invention is further illustrated by the following specific examples which are not intended in any way to limit the scope of the invention.

Examples

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Infection of NHBE cells by adenovirus serotypes of subgroup D Example 1 Normal human bronchial epithelial ("NHBE") cells were obtained from Clonetics (San Diego, CA), and plated on Costar (Cambridge, MA) Transwell-Clear polyester membranes that were pre-coated with human placental collagen. The wells were placed in a cluster plate and cells were fed every day for one week by changing the medium in both the well and the plate. After one week the media was removed from the wells to create an air-liquid interface, and the cells were then fed only by changing the medium in the cluster plate, every other day for one week. Cells were infected at an moi of 1 by adding virus (see below) to the transwell, followed by an incubation time of 1.5-2 hours. At the end of the incubation period, the medium was removed and the cells were gently rinsed with fresh medium. Thirty-six hours postinfection the cells were fixed with 1:1 acetone:methanol, permeablized with a solution of 0.05% Tween 20 in PBS, and stained with FITC labeled anti-hexon antibody (Chemicon, Temecula, CA) to visualize cells that had been productively infected (i.e. to visualize virus replication). Cells were also subjected to the DAPI staining procedure in order to visualize the total number of nuclei. The results could be readily determined upon simple inspection.

Wild type Ad serotypes within subgroup D that were tested included 9, 15, 17, 19, 20, 22, 26, 27, 28, 30, and 39 (all from the American Type Culture Collection, Rockville, MD). An Ad 2 (obtained as DNA from BRL, Gaithersburg, MD, and used to transfect 293 cells in order to generate virus stock) was used as a control. Infection observed with all of the subgroup D serotypes was superior to that observed with Ad 2, with the best results being achieved with Ad 9, Ad 17, Ad 20, Ad 22, and Ad 30.

Additionally, it was determined that each of the above-mentioned serotypes of subgroup D was more effective in the NHBE cell assay under similar circumstances than any other serotype tested than belongs to a subgroup other than D. In this regard, the following serotypes were also tested: 31(subgroup A); 3(subgroup B); 7(subgroup B); 7a(subgroup B); 4(subgroup E); and 41(subgroup F). In a further experiment, serotype 35 (subgroup A) may have performed as well as the least effective members of subgroup D that were tested.

Example 2 Infection of clinical isolate bronchial epithelial cells

Following generally the procedures of Example 1, human bronchial epithelial cells recovered from healthy human volunteers were infected with either Ad 2 (as above, Ad 2 DNA was obtained from BRL, and this DNA was used to transfect 293 cells to generate virus) (Figure 1), or Ad 17 (from ATCC) (Figure 2), all at an moi of 50. Cells were left in contact with virus for 30 minutes, 3 hours, or 12 hours.

The increased tropism of Ad 17 for human bronchial epithelial cells, compared with Ad 2, is readily apparent upon inspection of Figures 1 and 2. In the Figures, the right hand columns (panels D, E, and F, stained in blue) show total numbers of cells present (from DAPI staining as above), whereas the left hand columns (panels A, B, and C, stained in green) quantify adenovirus hexon protein present in the infected cells (from FITC-labeled anti-hexon anitbody, as above). Panels A and D result from 30 minute incubation times, panels B and E result from 3 hour incubation times, and panels C and F result from 12 hour incubation times. As measured by the technique employed, infection of airway epithelia by Ad 17 is at least 50 fold greater than by Ad 2 for the thirty minute incubation time.

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Example 3 Binding of Ad 2 and Ad 17 to human nasal polyp cell isolates 293 cells, a complementing cell line developed by Graham et al. (see Gen. Virol., 36, 1977, pp. 59-72), were infected with either wild type Ad 2 or wild type Ad 17. Five hours post-infection the media was removed and replaced with methionine

free media containing S³⁵ metabolic label (Amersham). After an additional six hours, fresh media was added and the labeling was allowed to proceed for a total of 18 hours, after which the S³⁵ media was removed and replaced with fresh media. Thirty hours post-infection the cells were harvested and lysed and the labeled Ad 2 or Ad 17 viruses were purified by CsCl gradient centrifugation. The recovered viruses were then used in an assay to determine their relative binding efficiency on human nasal polyp cells.

In order to perform the assay, ciliated human airway epitehlial cells were recovered from nasal polyps of healthy volunteers. The results from two such isolates, NP-14 and NP-15, are reported here (see Figure 3). Radiolabeled virus was then incubated with the isolated cells in wells for specified times (5 or 30 minutes, see Figure 3). The cells were then rinsed and measured for radioactivity. Binding as reported in Figure 3 indicates the percent of input radioactivity that is cell associated. It was determined that for both cell isolate populations, using either 5 or 30 minute incubations, cell associated radioactivity was 10-fold enhanced if Ad 17 rather than Ad 2 was used.

Example 4 Fiber competition

20 A549 cells (a human lung carcinoma line, obtained from the American Type Culture Collection as ATCC CCL-185) were plated at 3 x 10⁴ cells per well in 96-well dishes. Since the number of receptor sites for adenovirus fiber on the cell surface has been estimated to be approximately 10⁵ receptors per cell, the receptors in the plated cells were saturated, in this example, with 0.1µg of purified full length Ad 2 fiber protein (obtained from Paul Freimuth, Brookhaven National Laboratory, Upton, NY), which corresponds to approximately 100 molecules of fiber per receptor. Cells were incubated with Ad 2 fiber in PBS for two hours at 37°C.

The cells were subsequently infected at an moi of 1 (using either Ad 2 provided as above, or wild type Ad 17) for one hour, after which the cells were rinsed, and fresh mediium was added. Control cultures were incubated with PBS with no added protein for two hours and then subsequently infected as described above. Forty hours post-infection the cells were fixed with 1:1 acetone:methanol, permeablized with 0.05% Tween 20 in PBS and stained with FITC labeled anti- Ad 2 hexon antibody, as described in Example 1. As determined by this assay, the number of cells infected (stained) with Ad 2 was reduced by approximately 90% in cultures that were pre-incubated with Ad 2 fiber as compared to control cultures. However, no effect on Ad 17 infection was observed by the pre-incubation of A549 cells with full length Ad 2 fiber.

Example 5 Use of Ad 2 fiber knob in a binding competition experiment with Ad 2

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Further competition experiments were performed with Ad 2 and Ad 17 fiber knobs that had been expressed and purified from E. coli. DNA sequences encoding both protein fragments were designed so that the fiber knobs expressed therefrom would contain histidine tags in order to permit nickel- column purification. The yield of soluble fiber knob trimer, purified by the Ni-NTA method (Qiagen, Chatsworth, CA) , was $\sim 25 \mu g/50 ml$ culture. A significant portion of the total knob protein expressed appeared to remain in a monomeric (and insoluble) form. The soluble trimeric material obtained was used for a preliminary competition experiment. Wild type Ad 2 and Ad 17 were used to infect A549 cells, or cells that had been preincubated with excess (about 100 molecules of trimer per receptor) Ad 2 fiber knob or Ad 17 fiber knob. The results indicated that Ad 2 fiber knob, but not Ad 17 knob, could block Ad 2 infection. Additionally, Ad 17 infection was not blocked by E. coliexpressed fiber knobs of either serotype, suggesting that the mechanism of Ad 2 and Ad 17 infections is different.

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Example 6 Construction of the chimeric vector Ad2/βgal-2/fiber Ad 17

The vector Ad2/βgal-2 was constructed as follows. A CMV§gal expression cassette was constructed in a pBR322-based plasmid that contained Ad 2 nucleotides 1-10,680 from which nucleotides 357-3328 were deleted. The deleted sequences were replaced with (reading from 5' to 3'): a cytomegalovirus immediate early promoter (obtained from pRC/CMV, Invitrogen), lacZ gene encoding §-galactosidase with a nuclear localization signal, and an SV40 polyadenylation signal (nucleotides 2533-2729). The resulting plasmid was used to generate Ad2/βgal-2 by recombination with Ad2E4ORF6 (D. Armentano et al., Human Gene Therapy , 6, 1995, pp 1343 -1353).

A chimeric Ad2/ β gal-2/fiber Ad 17 viral vector (Figure 4) was then contructed as follows. pAdORF6 (D. Armentano et al., Human Gene Therapy , 6, 1995, pp 1343 -1353 was cut with Nde and BamHI to remove Ad 2 fiber coding and polyadenylation signal sequences (nucleotides 20624-32815). An NdeI-BamHI fragment containing Ad 17 fiber coding sequence (nucleotides 30984-32095) was generated by PCR and ligated along with an SV40 polyadenylation signal into NdeI-BamHI cut pAdORF6 to generate pAdORF6fiber17. This plasmid was cut with PacI and then ligated to PacI-cut Ad2/ β gal-2 DNA to generate Ad2/ β gal-2 fiber 17. Any desired transgene may be substituted in this construct for the reporter gene.

A similar construct can be prepared using a DNA sequence that encodes Ad 17 penton base instead of Ad 17 fiber. Alternatively, only a subregion of the penton base of Ad 2 need be subject to replacement, such as by inserting into the vector a nucleotide encoding sequence corresponding to any amino acid subsequence of Ad 17 penton base amino acids 283-348 (see the marked sequence in Figure 5A) in replacement for any subsequence of Ad 2 penton base amino acids 290-403. Preferrably, the replaced sequence of Ad 2 and the inserted sequence of Ad 17 includes the RGD domain of each. Use of nucleotide sequence corresponding to penton base amino acid sequence for other subgroup D serotypes is also within the

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practice of the invention. It is also within the scope of the invention to replace a subregion of the fiber protein in the Ad 2 vector with a subregion from another adenovirus scrotype, for example, Ad 17.

Example 7 Ad2/βgal-2f17 shows increased infection efficiency on human airway explants

Both human and monkey trachea explants, about 1 cm², were placed on top of an agar support. Each explant was infected at an moi of 200 of either Ad2/βgal-2 or Ad2/βgal-2f17 assuming a cell density of 1 x 10⁶ per cm² of explant. Explants were exposed to virus for three hours and were then rinsed with NHBE media. Two days post-infection explants were stained with X-gal and infection efficiency was assessed. On the monkey explants Ad2/βgal-2 gave rise to a higher infection efficiency than Ad2/βgal-2f17. Patches of stained cells were detected in explants exposed to Ad2/βgal-2f17. A different result was obtained on human trachea explants. On these explants Ad2/βgal-2f17 infection gave rise to a much higher infection efficiency than Ad2/βgal-2f17 stained with X-gal whereas very few cells were stained in explants exposed to Ad2/βgal-2f17 stained with X-gal whereas very few cells were stained in explants exposed to Ad2/βgal-2. No background staining was observed in either monkey or human explants that were not exposed to virus.

The results indicate that the exchange of Ad 2 fiber for Ad 17 fiber in Ad2/ β gal-2f17 was suffficient to significantly increase infection efficiency of human tracheal airway cells by an adenovirus type 2 based vector.

25 Example 8 Adenovirus subgroup screening on human cancer cell lines

Identification of adenovirus subgroup that best infects a particular tumor type may be useful in designing vectors to optimally target cancer cells in vivo. In order to determine the adenovirus subgroup that best infects a particular type of cancer cell, cancer cells were seeded into a 96 well plate and infected with and moi of 5. Infection

efficiency was determined by staining of infected cells using an anti-hexon antibody. The adenovirus subgroups were represented by the following serotypes: A: Ad 31; B: Ad 3; C: Ad 2; D: Ad 17; E: Ad 4; and F: Ad 41.

Subgroup D (Ad 17) has a significantly higher infection rate of the colon
cancer cell line CaCo-2 than other cell types, with an infection rate of 70%, while Ad
2 only infected 20% of the cells (Figure 9).

Subgroup D (Ad 17) was effective in infecting ovarian cancer cell line SK-OV3. Infection was measured at 90% (FIgure 10).

10 Sequence Listing

Included herewith on the following pages are informal copies of SEQ ID NO: 1 through SEQ ID NO: 3.

		-	.0	
	CATCATCAAT			
	TTTAGGGCGG			
	ACGGCTAACG			
	CGTATAAAAA			
	TATGAGGTAA			
	GAGGAAGTGA			
	AGAGACTTTG			
	TCCGTGTCAA			
	GTCGAGCCCG			
	CTCCCAGAGT			
	CATGGCCGCA			
	AACTCCGTTC			
	GGAGGACGAC			
	${\tt GGCTGACATA}$			
	ACCTGAATTG			
	CAGCGATTCA			
	TGTGGTTGTG			
	ATCCTGCCAG			
	CATGAAAAAG			
	GCTTAAGACA			
	${\tt TAGGTCCGGT}$			
	${\tt TCTGTCAGGC}$			
	GAGGCGAGCA			
	TTTGGACCTG			
	${\tt AATAAAGTTG}$			
	$\tt GGCGGGGCTT$			
	AGTTCCTGAT			
	AGGATAGTTC			
	GCCTGGTGTA			
	GCTCTGGCCT			
	TCCACAGCCT			
	TTCTGGTTGA			
	CAGCCATGCA			
	GGCTTCTACA			
	AGGAAGAAAT			
	AAGAGGAGTT			
	ATCCATGGCC			
	CGAGCTGACG			
	ACAGCAGGAG			
	AAAAACCCAT GATAGCCCTG			
	TGCTACATCT			
	AGGTGTTGCA			
	AACATGAAGT			
	ACCCTGCATG			
	TCCAAGATCA			
	AGCGAGATGT			
	GGCAATGCTA			
	GGCACAGCCT			
	ATGCTGACTG			
	CCAGAAAGAA			
	GCGCCAGAAG			
	TGGAGAACGA			
	TGTACAAGAT			
	GCAGACACAC			
	ACCTGGTGAT			
	GGTAGGTTTG			
1		 	-	

3421 CTTTTGCTT TTCTGCAGAC ATCATGAACG GGACCGGCGG GGCCTTCGAA GGGGGGCTTT 3481 TTAGCCCTTA TTTGACAACC CGCCTGCCAG GATGGGCCGG AGTTCGTCAG AATGTGATGG 3541 GATCGACGGT GGACGGCCC CCAGTGCTTC CAGCAAATTC CTCGACCATG ACCTACGCGA 3601 CCGTGGGGAA CTCGTCGCTT GACAGCACCG CCGCAGCCGC GGCAGCCGCA GCCGCCATGA 3661 CAGCGACGAG ACTGGCCTCG AGCTACATGC CCAGCAGCAG CAGTAGCCCC TCTGTGCCCA 3721 GTTCCATCAT CGCCGAGGAG AACTGCTGGC CCTGCTGGCC GAGCTGGAAG CCCTGAGCCG 3781 CCAGCTGGCC GCCTGACCC AGCAGGTGTC CGAGCTCCGC GAACAGCAGC AGCAAAATAA 3841 ATGATTCAAT AAACACATAT TCTGATTCAA ACAGCAAAGC ATCTTTATTA TTTATTTTTT 3901 CGCGCGCGT AGGCCCTGGT CCACCTCTCC CGATCATTGA GAGTGCGGTG GATTTTTTCC 3961 AAGACCCGGT AGAGGTGGGA TTGGATGTTG AGGTACATGG GCATGAGCCC GTCCCGGGGG 4021 TGGAGGTAGC ACCACTGCAT GGCCTCGTGC TCTGGGGTCG TGTTGTAGAT GATCCAGTCA 4081 TAGCAGGGC GCTGGGCGTG GTGCTGGATG ATGTCCTTGA GGAGGAGACT GATGGCCACG 4141 GGGAGCCCCT TGGTGTAGGT GTTGGCAAAG CGGTTGAGCT GGGAGGGATG CATGCGGGGG 4201 GAGATGATGT GCAGTTTGGC CTGGATCTTG AGGTTGGCGA TGTTGCCACC CAGATCCCGC 4261 CGGGGGTTCA TGTTGTGCAG GACCACCAGG ACGGTGTAGC CCGTGCACTT GGGGAACTTA 4321 TCATGCAACT TGGAAGGGAA TGCGTGGAAG AATTTGGAGA CGCCCTTGTG CCCGCCCAGG 4381 TTTTCCATGC ACTCATCCAT GATGATGGCG ATGGGCCCGT GGGCTGCGGC TTTGGCAAAG 4441 ACGTTTCTGG GGTCAGAGAC ATCATAATTA TGCTCCTGGG TGAGATCATC ATAAGACATT 4501 TTAATGAATT TTGGGCGGAG GGTGCCAGAT TGGGGGACGA TGGTTTCCCT CGGGCCCCGG 4561 GGCGAAGTTC CCCTCGCAGA TCTGCATCTC CCAGGCTTTC ATCTCGGAGG GGGGGATCAT 4621 GTCCACCTGC GGGGCGATGA AAAAAACGGT TTCCGGGGCG GGGGTGATGA GCTGCGAGGA 4681 GAGCAGGTTT CTCAACAGCT GGGACTTGCC GCACCCGGTC GGGCCGTAGA TGACCCCGAT 4741 GACGGGTTGC AGGTGGTAGT TCAAGGACAT GCAGCTGCCG TCGTCCCGGA GGAGGGGGGC 4801 CACCTCGTTG AGCATGTCTC TAACTTGGAG GTTTTCCCGG ACGAGCTCGC CGAGGAGGCG 4861 GTCCCCGCCC AGCGAGAGGA GCTCTTGCAG GGAAGCAAAG TTTTTCAGGG GCTTGAGTCC 4921 GTCGGCCATG GGCATCTTGG CGAGGGTCTG CGAGAGGAGT TCGAGACGTC CCAGAGCTCG 4981 GTGACGTGCT CTACGGCATC TCGATCCAGC AGACTTCCTC GTTTCGGGGG TTGGGACGAC 5041 TGCGACTGTA GGGCACGAGA CGATGGGCGT CCAGCGCGGC CAGCGTCATG TCCTTCCAGG 5101 GTCTCAGGGT CCGCGTGAGG GTGGTCTCCG TCACGGTGAA GGGGTGGGCC CCTGGCTGGG 5161 CGCTTGCAAG GGTGCGCTTG AGACTCATCC TGCTGGTGCT GAAACGGGCA CGGTCTTCGC 5221 CCTGCGCGTC GGCGAGATAG CAGTTGACCA TGAGCTCGTA GTTGAGGGCC TCGGCGGCGT 5281 GGCCCTTGGC GCGGAGCTTG CCCTTGGAAG AGCGTCCGCA GGCGGGACAG AGGAGGGATT 5401 AGTGGGCGCA GACGGTCTCG CACTCGACGA GCCAGGTGAG CTCGGGCTGC TCGGGGTCAA 5461 AAACCAGTTT TCCCCCGTTC TTTTTGATGC GCTTCTTACC TCGCGTCTCC ATGAGTCTGT 5521 GTCCGCGCTC GGTGACAAAC AGGCTGTCGG TGTCCCCGTA GACGGACTTG ATTGGCCTGT 5581 CCTGCAGGGG CGTCCCGCGG TCCTCCTCGT AGAGAAACTC GGACCACTCT GAGACAAAGG 5641 CGCGCGTCCA CGCCAAGACA AAGGAGGCCA CGTGCGAGGG GTAGCGGTCG TTGTCCACCA* 5701 GGGGGTCCAC CTTTTCCACC GTGTGCAGAC ACATGTCCCC TTCCTCCGCA TCCAAGAAGG 5761 TGATTGGCTT GTAGGTGTAG GCCACGTGAC CAGGGGTCCC CGACGGGGG GTATAAAAGG 5821 GGGCGGGTCT GTGCTCGTCC TCACTCTCTT CCGCGTCGCT GTCCACGAGC GCCAGCTGTT 5881 GGGGTAGGTA TTCCCTCTCG AGAGCGGGCA TGACCTCGGC ACTCAGGTTG TCAGTTTCTA 5941 GAAACGAGGA GGATTTGATG TTGGCTTGCC CTGCCGCAAT GCTTTTTAGG AGACTTTCAT 6001 CCATCTGGTC AGAAAAGACT ATTTTTTTAT TGTCAAGCTT GGTGGCAAAG GAGCCATAGA 6061 GGGCGTTGGA GAGAAGCTTG GCGATGGATC TCATGGTCTG ATTTTTGTCA CGGTCGGCGC 6121 GCTCCTTGGC CGCGATGTTG AGCTGGACAT ATTCGCGCGC GACACACTTC CATTCGGGAA 6181 AGACGGTGGT GCGCTCGTCG GGCACGATCC TGACGCGCCA GCCGCGGTTA TGCAGGGTGA 6241 CCAGGTCCAC GCTGGTGGCC ACCTCGCCGC GCAGGGGCTC GTTAGTCCAG CAGAGTCTGC 6301 CGCCCTTGCG CGAGCAGAAC GGGGGCAGCA CATCAAGCAG ATGCTCGTCA GGGGGGTCCG 6361 CATCGATGGT GAAGATGCCG GGACAGAGTT TCTTGTCAAA ATAGTCTATT TTTGAGGATG 6421 CATCATCCAA GGCCATCTGC CACTCGCGGG CGGCCATTGC TCGCTCGTAG GGGTTGAGGG 6481 GCGGACCCCA CGGCATGGGA TGCGTGAGGG CGGAGGCGTA CATGCCGCAA ATGTCGTAAA 6541 CATAGATGGG CTCCGAGAAG ATGCCGATGT TGGTGGGATA ACAGCGCCCC CCGCGGATGC 6601 TGGCGCGCAC GTATTCATAC AACTCGTGCG AGGGGCCAAG AAGGCCGGGG CCGAAATTGG 6661 TGCGCTGGGG CTGCTCGGCG CGGAAAACAA TCTGGCGAAA GATGGCGTGC GAGTTGGAGG 6721 AGATGGTGGG CCGTTGGAAG ATGTTAAAGT GGGCGTGGGG CAAGCGGACC GAGTCGCGGA 6781 TGAAGTGCGC GTAGGAGTCT TGCAGCTTGG CGACGAACTC GGCGGTGACG AGAACGTCCA

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6841 TGGCGCAGTA GTCCAGCGTT TCGCGGATGA TGTCATAACC CGCCTCTCCT TTCTTCTCCC 6901 ACAGCTCGCG GTTGAGGGCG TATTCCTCGT CATCCTTCCA GTACTCCCGG AGCGGGAATC 6961 CTCGATCGTC CGCACGGTAA GAGCCCAGCA TGTAGAAATG GTTCACGGCC TTGTAGGGAC 7021 AGCAGCCCTT CTCCACGGG AGGGCGTAAG CTTGTGCGGC CTTGCGGAGC GAGGTGTGCG 7081 TCAGGGCGAA GGTGTCCCTG ACCATGACTT TCAAGAACTG GTACTTGAAA TCCGAGTCGT 7141 CGCAGCCGCC GTGCTCCCAT AGCTCGAAAT CGGTGCGCTT CTTCGAGAGG GGGTTAGGCA 7201 GAGCGAAAGT GACGTCATTG AAGAGAATCT TGCCTGCTCG CGGCATGAAA TTGCGGGTGA 7261 TGCGGAAAGG GCCCGGGACG GAGGCTCGGT TGTTGATGAC CTGGGCGGCG AGGACGATCT 7321 CGTCGAAGCC GTTGATGTTG TGCCCGACGA TGTAGAGTTC CATGAATCGC GGGCGGCCTT 7381 TGATGTGCGG CAGCTTTTTG AGCTCCTCGT AGGTGAGGTC CTCGGGGCAT TGCAGGCCGT 7441 GCTGCTCGAG CGCCCATTCC TGGAGATGTG GGTTGGCTTG CATGAAGGAA GCCCAGAGCT 7501 CGCGGGCCAT GAGGGTCTGG AGCTCGTCGC GAAAGAGGCG GAACTGCTGG CCCACGGCCA 7561 TCTTTTCGGG TGTGACGCAG TAGAAGGTGA GGGGGTCCCG CTCCCAGCGA TCCCAGCGTA 7621 AGCGCGCGC TAGATCGCGA GCAAGGGCGA CCAGCTCTGG GTCCCCCGAG AATTTCATGA 7681 CCAGCATGAA GGGGACGAGC TGCTTGCCGA AGGACCCCAT CCAGGTGTAG GTTTCTACAT 7741 CGTAGGTGAC AAAGAGCCGC TCCGTGCGAG GATGAGAGCC GATTGGGAAG AACTGGATTT 7801 CCTGCCACCA GTTGGACGAG TGGCTGTTGA TGTGATGAAA GTAGAAATCC CGCCGGCGAA 7861 CCGAGCACTC GTGCTGATGC TTGTAAAAGC GTCCGCAGTA CTCGCAGCGC TGCACGGGCT 7921 GTACCTCATC CACGAGATAC ACAGCGCGTC CCTTGAGGAG GAACTTCAGG AGTGGCGGCC 7981 CTGGCTGGTG GTTTTCATGT TCGCCTGCGT GGGACTCACC CTGGGGCTCC TCGAGGACGG 8041 AGAGGCTGAC GAGCCCGCGC GGGAGCCAGG TCCAGATCTC GGCGCGGCGG GGGCGGAGAG 8101 CGAAGACGAG GGCGCGCAGT TGGGAGCTGT CCATGGTGTC GCGGAGATCC AGGTCCGGGG 8161 GCAGGGTTCT GAGGTTGACC TCGTAGAGGC GGGTGAGGGC GTGCTTGAGA TGCAGATGGT 8221 ACTTGATTTC TACGGGTGAG TTGGTGGCCG TGTCCACGCA TTGCATGAGC CCGTAGCTGC 8281 GCGGGGCCAC GACCGTGCCG CGGTGCGCTT TTAGAAGCGG TGTCGCGGAC GCGCTCCCGG 8341 CGGCAGCGGC GGTTCCGGCC CCGCGGGCAG GGGCGGCAGA GGCACGTCGG CGTGGCGCTC 8401 GGGCAGGTCC CGGTGTTGCG CCCTGAGAGC GCTGGCGTGC GCGACGACGC GGCGGTTGAC 8461 ATCCTGGATC TGCCGCCTCT GCGTGAAGAC CACTGGCCCC GTGACTTTGA ACCTGAAAGA 8521 CAGTTCAACA GAATCAATCT CGGCGTCATT GACGGCGGGC TGACGCAGGA TCTCTTGCAC 8581 GTCGCCCGAG TTGTCCTGGT AGGCGATCTC GGACATGAAC TGCTCGATCT CCTCCTCCTG 8641 GAGATCGCCG CGACCCGCGC GCTCCACGGT GGCGGCGAGG TCATTCGAGA TGCGACCCAT 8701 GAGCTGCGAG AAGGCGCCCA GGCCGCTCTC GTTCCAGACG CGGCTGTAGA CCACGTCCCC 8761 GTCGGCGTCG CGCGCGCGCA TGACCACCTG CGCGAGGTTG AGCTCCACGT GCCGCGCGAA 8821 GACGCCTAG TTGCGCAGGC GCTGGAAGAG GTAGTTGAGG GTGGTGGCGA TGTGCTCGGT 8881 GACGAAGAAG TACATGATCC AGCGGCGCAG GGGCATCTCG CTGATGTCGC CGATGGCCTC 8941 CAGCCTTTCC ATGGCCTCGT AGAAATCCAC GGCGAAGTTG AAAAACTGGG CGTTGCGGGC 9001 CGAGACCGTG AGCTCGTCTT CCAGGAGCCT GATGAGCTCG GCGATGGTGG CGCGCACCTC 9061 GCGCTCGAAA TCCCCGGGGG CCTCGTCCTC TTCCTCTTCT TCCATGACAA CCTCTTCTAT 9121 TTCTTCCTCT GGGGGCGGTG GTGGTGGCGG GGCCCGACGA CGACGGCGAC GCACCGGGAG 9181 ACGGTCGACG AAGCGCTCGA TCATCTCCCC GCGGCGGCGA CGCATGGTTT CGGTGACGGC 9241 GCGACCCCGT TCGCGAGGAC GCAGCGTGAA GACGCCGCCG GTCATCTCCC GGTAATGGGG 9301 CGGGTCCCCG TTGGGCAGCG AGAGGGCGCT GACGATGCAT CTTATCAATT GCGGTGTAGG 9361 GGACGTGAGC GCGTCGAGAT CGACCGGATC GGAGAATCTT TCGAGGAAAG CGTCTAGCCA 9421 ATCGCAGTCG CAAGGTAAGC TCAAACACGT AGCAGCCCTG TGGACGCTGT TAGAATTGCG 9481 GTTGCTAATG ATGTAATTGA AGTAGGCGTT TTTGAGGCGG CGGATGGTGG CGAGGAGGAC 9541 CAGGTCCTTG GGTCCCGCTT GCTGGATGCG GAGCCGCTCG GCCATGCCCC AGGCCTGGCC 9601 CTGACACCGG CTTAGGTTCT TGTAGTAGTC ATGCATGAGC CTCTCGATGT CATCACTGGC 9661 GGAGGCGGAG TCTTCCATGC GGGTGACCCC GACGCCCCTG AGCGGCTGCA CGAGCGCCAG 9721 GTCGGCGACG ACGCGCTCGG CGAGGATGGC CTGTTGCACG CGGGTGAGGG TGTCCTGGAA 9781 GTCGTCCATG TCGACGAAGC GGTGGTAGGC CCCTGTGTTG ATGGTGTAAG TGCAGTTGGC 9841 CATGAGCGAC CAGTTGACGG TCTGCAGGCC GGGCTGCACG ACCTCGGAGT ACCTGAGCCG 9901 CGAGAAGGCG CGCGAGTCGA AGACGTAGTC GTTGCAGGTG CGCACAAGGT ACTGGTATCC 9961 GACTAGGAAG TGCGGCGGCG GCTGGCGGTA GAGCGGCCAG CGCTGGGTGG CCGGCGCGCC 10021 CGGGGCCAGG TCCTCGAGCA TGAGGCGGTG GTAGCCGTAG AGGTAGCGGG ACATCCAGGT 10081 GATGCCGGCA GCGGTGGTGG AGGCGCGCGG GAACTCGCGG ACGCGGTTCC AGATGTTGCG 10141 CAGCGGCAGG AAATAGTCCA TGGTCGGCAC GGTCTGGCCG GTGAGACGCG CGCAGTCATT 10201 GACGCTCTAG AGGCAAAAAC GAAAGCGGTT GAGCGGGCTC TTCCTCCGTA GCCTGGCGGA

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10261	ACGCAAACGG	GTTAGGCCGC	GCGTGTACCC	CGGTTCGAGT	CCCCTCGAAT	CAGGCTGGAG
10321	CCGCGACTAA	CGTGGTATTG	GCACTCCCGT	CTCGACCCGA	GCCCGATAGC	CGCCAGGATA
10381	CGCGGGAAGA	GCCCTTTTTG	CCGGCCGARG	GGAGTCGCTA	GACTTGAAAG	CGGCCGAAAA
10441	CCCCGCCGGG	TAGTGGCTCG	CGCCCGTAGT	CTGGAGAAGC	ATCGCCAGGG	TTGAGTCGCG
10501	GCAGAACCCG	GTTCGCGGAC	GGCCGCGGCG	AGCGGGACTT	GGTCACCCCG	CCGATTTAAA
10561	GACCCACAGC	CAGCCGACTT	CTCCAGTTAC	GGGAGCGAGC	CCCCTTTTTT	CTTTTTGCCA
10621	GATGCATCCC	GTCCTGCGCC	AAATGCGTCC	CACCCCCCCG	GCGACCACCG	CGACCGCGGC
10681	CGTAGCAGGC	GCCGGCGCTA	GCCAGCCACA	GCCACAGACA	GAGATGGACT	TGGAAGAGGG
10741	CGAAGGGCTG	GCGAGACTGG	GGGCGCCTTC	CCCGGAGCGA	CACCCCCGCG	TGCAGCTGCA
10801	GAAGGACGTG	CGCCCGGCGT	ACGTGCCTGC	GCAAAACCTG	TTCAGGGACC	GCAGCGGGGA
10861	GGAGCCCGAG	GAGATGCGCG	ACTGCCGGTT	TCGGGCGGGC	AGGGAGCTGC	GCGAGGGCCT
10921	GGACCGCCAG	CGCGTGCTGC	GCGACGAGGA	TTTCGAGCCG	AACGAGCAGA	CGGGGATCAG
10981	CCCCGCGCGC	GCGCACGTGG	CGGCGGCCAA	CCTGGTGACG	GCCTACGAGC	AGACGGTGAA
		AACTTCCAAA				
11101	GGAGGTGGCC	CTGGGCCTGA	TGCACCTGTG	GGACCTGGCG	GAGGCCATCG	TGCAGAACCC
11161	GGACAGCAAG	CCTCTGACGG	CGCAGCTGTT	CCTGGTGGTA	CAGCACAGCA	GGGACAACGA
11221	GGCGTTCAGG	GAGGCGCTGC	TAAACATCGC	CGAGCCCGAG	GGTCGCTGGC	TGCTGGAGCT
		TTGCAGAGCA				_
		AACTACTCGG				
		GTGCCCATAG				
		CTGACGCTGA				
		GCGAGCCGGC				
		GTAGGGGGCG				
		CAGCCGAGCC				
		GAAGAGGAGG				
		GCAGCAAGCC				
	-	ATCGGACGAC				
		GTCCTTTAGA				
		TTCTCGGACC				
		CAAGGCCATC				
		CCGCTACAAC				
		AGCCGTGGCG				
		CGCCTTCCTG				
		TATCAGCGCG				
		CCCGGACTAC				
		TTTCAAGAAC				
		GAGCAGCTTG				
		CAGTGGCAGC				
		CATAGGCCAG				
		GCTGGGGCAG				
		ACAGCAGAAG				
		TGTGCAGCAG				
		GGACATGACC				
		TAAGCTAATG				
		CATTTTGAAC				
		CGACCCCAAC				
		GCAAAAGCGC				
'		CTTTCCTAGC				
		CCGGCCGCGC				
		GGTCAAGAAC				
		GAAGACCTAC				
		CCGGCAGCGG				
		CTTGGGCGGG				
		ACGGATGTTT				
		TTAGAGATGA				
		GCGCAGGCGA				
		AGAAACAGCA				
17021	TUCOCHOCOC	NONMACAGEA	TICGIIMCIC	GOVOC 1 GGC 1	CCGIIGIACG	ACACCACICG

13681 CGTGTACTTG GTGGACAACA AGTCGGCGGA CATCGCTTCC CTGAACTATC AAAACGACCA 13741 CAGCAACTTC CTGACCACGG TGGTGCAGAA CAACGATTTC ACCCCCGCCG AGGCTAGCAC 13801 GCAGACGATA AATTTTGACG AGCGGTCGCG GTGGGGCGGT GATCTGAAGA CCATTCTGCA 13861 CACCAACATG CCCAATGTGA ACGAGTACAT GTTCACCAGC AAGTTTAAGG CGCGGGTGAT 13921 GGTGGCTAGA AAACACCCAC AGGGGGTAGA AGCAACAGAT TTAAGCAAGG ATATCTTAGA 13981 GTATGAGTGG TTTGAGTTTA CCCTGCCCGA GGGCAACTTT TCCGAGACCA TGACCATAGA 14041 CCTGATGAAC AACGCCATCT TGGAAAACTA CTTGCAAGTG GGGCGGCAAA ATGGCGTGCT 14101 GGAGAGCGAT ATTGGAGTCA AGTTTGACAG CAGAAATTTC AAGCTGGGCT GGGACCCTGT 14161 GACCAAGCTG GTGATGCCAG GGGTCTACAC CTACGAGGCC TTTCACCCGG ACGTGGTGCT 14221 GCTGCCGGGC TGCGGGGTGG ACTTCACAGA GAGCCGCCTG AGCAACCTCC TGGGCATTCG 14281 CAAGAAGCAA CCTTTCCAAG AGGGCTTCAG AATCATGTAT GAGGATCTAG AAGGGGGCAA 14341 CATCCCCGCC CTGCTGGATG TGCCCAAGTA CTTGGAAAGC AAGAAGAAGT TAGAGGAGGC 14401 ATTGGAGAAT GCTGCTAAAG CTAATGGTCC TGCAAGAGGA GACAGTAGCG TCTCAAGAGA 14461 GGTTGAAAAG GCAGCTGAAA AAGAACTTGT TATTGAGCCC ATCAAGCAAG ATGATACCAA 14521 GAGAAGTTAC AACCTCATCG AGGGAACCAT GGACACGCTG TACCGCAGCT GGTACCTGTC 14581 CTATACCTAC CGGGACCCTG AGAACGGGGT GCAGTCGTGG ACGCTGCTCA CCACCCCGGA 14641 CGTCACCTGC GGCGCGGAGC AAGTCTACTG GTCGCTGCCG GACCTCATGC AAGACCCCGT 14701 CACCTTCCGT TCTACCCAGC AAGTCAGCAA CTACCCCGTG GTCGGCGCCG AGCTCATGCC 14761 CTTCCGCGCC AAGAGCTTTT ACAACGACCT CGCCGTCTAC TCCCAGCTCA TCCGCAGCTA 14821 CACCTCCCTC ACCCACGTCT TCAACCGCTT CCCCGACAAC CAGATCCTCT GCCGTCCGCC 14881 CGCGCCCACC ATCACCACCG TCAGTGAAAA CGTGCCTGCT CTCACAGATC ACGGGACGCT 14941 ACCGCTGCGC AGCAGTATCC GCGGAGTCCA GCGAGTGACC GTCACTGACG CCCGTCGCCG 15001 CACCTGTCCC TACGTCTACA AGGCCCTGGG CATAGTCGCG CCGCGTGTGC TTTCCAGTCG 15061 CACCTTCTAA AAAATGTCTA TTCTCATCTC GCCCAGCAAT AACACCGGCT GGGGTATTAC 15121 TAGGCCCAGC AGCATGTACG GAGGAGCCAA GAAACGTCCC AGCAGCACCC CGTCCGCGTC 15181 CGCGGCCACT TCCGCGCTCC GTGGGGCGCT TACAAGCGCG GGCGGACTGC CACCGCCGCC 15241 GCCGTGCGCA CCACCGTCGA CGACGTCATC GACTCGGTGG TCGCCGACGC GCGCAACTAT 15301 ACTCCCGCCC CTTCGACCGT GGACGCGGTT CATTGACAGC GTGGTGGCGA CGCGGCGGCG 15361 ATATGCCAGA CGCAAGAGCC GGCGGCGGA CGGATCGCCC AGGCGCCATT CGGAGCACGC 15421 CCGCCATGGG GCGCCGCCC AGCTCTGCTG CGCCGCGCCA GACGCACGGG CCGCCGGGCC 15481 ATGATGCGAG CCGCGCGCCG CGCCGCCACT GCACCCCCG CAGGCAGGAC TCGCAGACGA 15541 GCGGCCGCCG CCGCCGCCGC GGCCATCTCT AGCATGACCA GACCCAGGCG CGGAAACGTG 15601 TACTGGTGC GCGACTCCGT CACGGGCGTG CGCGTGCCCG TGCGCACCCG TCCTCCTCGT 15661 CCCTGATCTA ATGCTTGTGT CCTCCCCGC AAGCGACGAT GTCAAAGCGC ATCTACAAGA 15721 GAGATGCTCC AGGTCGTCGC CCCGGAGATT TACGGACCAC CCCAGGCGGA CCAGAAACCC 15781 CGCAAAATCA AGCGGGTTAA AAAAAAGGAT GAGGTGGACG AGGGGGCAGT AGAGTTTGTG 15841 CGCGAGTTCG CTCCGCGGCG GCGCGTAAAT TGGAAGGGGC GCAGGTGCAC GCGTGTTGCG 15901 GCCCGGCACG GCGGTGGTGT TCACGCCCGG CGAGCGGTCC TCGGTCAGGA GCAAGCGTAG 15961 CTATGACGAG GTGTACGGCG ACGACGACAT CCTGGACCAG GCGGCAGAGC GGGCGGGCGA 16021 GTTTGCCTAC GGGAAGCGGT CGCGCGAAGA GGAGCTGATC TCGCTGCCGC TGGACGAGAG 16081 CAATCCCACG CCGAGCCTGA AGCCCGTGAC CTGCAGCAGG TGCTGCCCCA GGCGGTGCTG 16141 CTGCCGAGCC GCGGGATCAA GCGCGAGGGC GAGAACATGT ACCCGACCAT GCAGATCATG 16201 GTGCCCAAGC GCCGCGCGT GGAGGAAGTG CTGGACACCG TGAAAATGGA TGTGGAGCCC 16261 GAGGTCAAGG TGCGCCCCAT CAAGCAGGTG GCGCCGGGCC TGGGCGTGCA GACCGTGGAC 16321 ATTCAGATCC CCACCGACAT GGATGTCGAC AAAAAACCCT CGACCAGCAT CGAGGTGCAG 16381 ACCGACCCT GGCTCCAGC CTCCACCGCT ACCGCTTCCA CTTCTACCGT CGCCACGGTC 16441 ACCGAGCCTC CCAGGAGGCG AAGATGGGGC CCCGCCAACC GGCTGATGCC CAACTACGTG 16501 TTGCATCCTT CCATTATCCC GACGCCGGCC TACCGCGGCA CCCGGTACTA CGCCAGCCGC 16561 AGGCGCCCAG CCAGCAAACG CCGCCGCCGC ACCGCCACCC GCCGCCGTCT GCCCCCCGCC 16621 CGCGTGCGCC GCGTAACCAA CGCGCCGGGG CCGCTCGCTC GTTCTGCCCA CCGTGCGCTA 16681 CCACCCCAGC ATCCTTTAAT CCGTGTGCTG TGATACTGTT GCAGAGAGAT GGCTCTCACT 16741 TGCCGCCTGC GCATCCCCGT TCCGAATTAC CGAGGAAGAT CCCGCCGCAG GAGAGGCATG 16801 GCAGGCAGCG GCCTGAACCG CCGCCGGCGG CGGCCCATGC GCAGGCGCCT GAGTGGCGGC 16861 TTTCTGCCCG CGCTCATCCC CATAATCGCG GCGCCATCG GCACGATCCC GGGCATAGCT 16921 TCCGTTGCGC TGCAGGCGTC GCAGCGCCGT TGATGTGCGA ATAAAGCCTC TTTAGACTCT 16981 GACACACCTG GTCCTGTATA TTTTTAGAAT GGAAGACATC AATTTTGCGT CCCTGGCTCC 17041 GCGCACGGC ACGCGGCCGT TCATGGGCAC CTGGAACGAG ATCGGCACCA GCCAGCTGAA

17101 CGGGGGCCC TTCAATTGGA GCAGTGTCTG GAGCGGGCTT AAAAATTTCG GCTCGACGCT 17161 CCGGACCTAT GGGAACAAGG CCTGGAATAG TAGCACGGGG CAGTTGTTGA GGGAAAAGCT 17221 CAAAGACCAG AACTTCCAGC AGAAGGTGGT GGACGGCCTG GCCTCGGGCA TTAACGGGGT 17281 GGTGGACATC GCGAACCAGG CAGTGCAGCG CGAGATAAAC AGCCGTCTGG ACCCGCGGCC 17341 GCCCACGGTG GTGGAGATGG AAGATGCAAC TCTTCCGCCG CCGAAGGGCG AGAAGCGGCC 17401 GCGGCCAGAT GCGGAGGAGA CGATCCTGCA GGTGGACGAG CCGCCTTCGT ACGAGGAGGC 17461 CGTGAAGGCC GGCATGCCCA CCACGCGCAT CATCGCGCCA CTGGCCACGG GTGTAATGAA 17521 ACCCGCCACC CTTGACCTGC CTCCACCACC CACGCCCGCT CCACCGAAGG CAGCTCCGGT 17581 TGTGCAGCCC CCTCCGGTGG CGACCGCCGT GCGCCGGTC CCCGCCCGCC GCCAGGCCCA 17641 GAACTGCCAG AGCACGCTGC ACAGTATTGT GGGCCTGGGA GTGAAAAGTC TGAAGCGCCG 17701 CCGATGCTAT TGAGAGAGAG GAAGGAGGAC ACTAAAGGGA GAGCTTAACT TGTATGTGCC 17761 TTACCGCCAG AGAACGCGCG AAGATGGCCA CCCCCTCGAT GATGCCGCAG TGGGCGTACA 17821 TGCACATCGC CGGGCAGGAC GCCTCGGAGT ACCTGAGCCC GGGTCTGGTG CAGTTTGCCC 17881 GCGCCACCGA CACGTACTTC AGCCTGGGCA ACAAGTTTAG GAACCCCACG GTGGCCCCGA 17941 CCCACGATGT GACCACGGAC CGGTCCCAGC GTCTGACGCT GCGCTTTGTG CCCGTGGATC 18001 GCGAGGACAC CAGTACTCGT ACAAGGCGCG CTTCACTCTG GCCGTGGGCG ACAACCGGGT 18061 GCTAGACATG GCCAGCACGT ACTTTGACAT CCGCGGCGTC CTGGACCGCG GTCCCAGTTT 18121 CAAACCCTAC TCGGGCACGG CTTACAACAG CCTTGCCCCC AAGGGCGCTC CCAATCCCAG 18181 TCAGTGGGTT GCCAAAGAAA ATGGTCAGGG AACTGATAAG ACACATACTT ATGGCTCAGC 18241 TGCCATGGGA GGAAGCAACA TCACCATTGA AGGTTTAGTA ATTGGAACTG ATGAAAAAGC 18301 TGAGGATGGC AAAAAAGATA TTTTTGCAAA TAAACTTTAT CAGCCAGAAC CTCAAGTAGG 18361 TGAAGAAAAC TGGCAAGAGT CTGAAGCCTT CTATGGAGGC AGAGCTCTTA AGAAAGACAC 18421 AAAAATGAAG CCCTGCTATG GCTCATTTGC AAGACCTACC AATGAAAAAG GCGGACAAGC 18481 TAAATTTAAG CCAGTGGAAG AGGGGCAGCA ACCTAAAGAT TATGACATAG ATTTGGCTTT 18541 CTTTGACACA CCTGGAGGCA CCATCACAGG AGGCACAGAC GAAGAATATA AAGCAGACAT 18601 TGTGTTGTAC ACTGAAAATG TCAACCTTGA AACCCCAGAC ACCCACGTGG TATACAAGCC 18661 AGGAAAAGAG GATGACAGTT CAGAAGTAAA TTTGACACAG CAGTCCATGC CCAACAGGCC 18721 TAACTACATT GGCTTCAGAG ACAACTTTGT GGGACTCATG TACTACAACA GTACTGGCAA 18781 CATGGGTGTG CTGGCTGGTC AGGCCTCTCA ATTGAATGCT GTGGTCGACT TGCAAGACAG 18841 AAACACCGAG CTGTCTTACC AGCTCTTGCT AGATTCTCTG GGTGACAGAA CCAGATACTT 18901 CAGCATGTGG AACTCTGCGG TGGATAGCTA TGATCCAGAT GTCAGGATCA TTGAAAATCA 18961 TGGTGTGGAA GATGAACTTC CAAACTATTG CTTCCCATTG AATGGCACTG GCACCAATTC 19021 AACATATCTT GGCGTAAAGG TGAAACCAGA TCAAGATGGT GATGTTGAAA GCGAGTGGGA 19081 TAAAGATGAT ACCATTGCAA GGCAGAATCA AATCGCCAAG GGCAACGTCT TTGCCATGGA 19141 GATCAACCTC CAGGCCAACC TGTGGAAGAG TTTTCTGTAC TCGAACGTGG CCTTGTACCT 19201 GCCCGACTCC TACAAGTACA CGCCGGCCAA TGTTACGCTG CCCGCCAACA CCAACACCTA 19261 CGAGTACATG AACGCCCGCG TGGTAGCCCC CTCGCTGGTG GACGCCTACA TCAACATAGG 19321 CGCCCGATGG TCGCTGGACC CCATGGACAA CGTCAACCCC TTCAACCACC ACCGCAATGC 19381 GGGCCTGCGC TACCGCTCCA TGCTTCTGGG CAACGGCCGC TACGTGCCCT TCCACATCCA 19441 AGTGCCCCAA AAGTTCTTTG CCATCAAGAA CCTGCTCCTG CTCCCGGGCT CCTACACCTA 19501 CGAGTGGAAC TTCCGCAAGG ATGTCAACAT GATCCTGCAG AGTTCCCTCG GCAACGACCT 19561 GCGCGTCGAC GGCGCCTCCG TCCGCTTCGA CAGCGTCAAC CTCTACGCCA CCTTCTTCCC 19621 CATGGCGCAC AACACCGCCT CCACCCTGGA AGCCATGCTG CGCAACGACA CCAACGACCA 19681 GTCCTTCAAC GACTACCTCT CGGCCGCCAA CATGCTCTAC CCCATCCCGG CCAAGGCCAC 19741 CAACGTGCCC ATCTCCATCC CCTCGCGCAA CTGGGCCGCT TTTCGCGGCT GGAGTTTCAC 19801 CCGTCTGAAA ACCAAGGAAA CTCCCTCCCT CGGCTCGGGT TTTGACCCCT ACTTTGTCTA 19861 CTCGGGCTCG ATCCCCTACC TTGACGGACC CTTTTACCTT AACCACACCT TCAAGAAAGT 19921 CTCCATCATG TTCGACTCCT CGGTCAGCTG GCCCGGCAAC GACCGGCTGC TCACGCCGAA 19981 CGAGTTCGAG ATCAAGCGCA GCGTCGACGG GGAAGGCTAC AACGTGGCCC AATGCAACAT 20041 GACCAAGGAC TGGTTCCTCG TCCAGATGCT CTCCCACTAC AACATCGGCT ACCAGGGCTT 20101 CCACGTGCCC GAGGGCTACA AGGACCGCAT GTACTCCTTC TTCCGCAACT TCCAGCCCAT 20161 GAGCAGGCAG GTGGTCGATG AGATCAACTA CAAGGACTAC AAGGCCGTCA CCCTGCCCTT 20221 CCAGCACAAC AACTCGGGCT TCACCGGCTA CCTTGCACCC ACCATGCGCC AAGGGCAGCC 20281 CTACCCGCC AACTTCCCCT ACCCGCTCAT CGGCCAGACA GCCGTGCCAT CCGTCACCCA 20341 GAAAAGTCTC CTCTGCGACA GGGTCATGTG GCGCATCCCC TTCTCCAGCA ACTTCATGTC 20401 CATGGGCGCC TTCACCGACC TGGGTCAGAA CATGTTCTAC GCCAACTCGG CCCACGCGCT 20461 CGACATGACC TTCGAGGTGG ACCCCATGGA TGAGCCCACC GTCCTCTATC TTCTCTTCGA

			- 3	4 -		
20521	AGTGTTCGAC	GTGGTCAGAG	TGCACCAGCC	GCACCGCGGC	GTCATCGAGG	CCGTCTACCT
20581	GCGCACGCCG	TTCTCCGCCG	GAAACGCCAC	CACCTAAGCA	TGAGCGGCTC	CAGCGAAAGA
20641	GAGCTCGCGT	CCATCGTGCG	CGACCTGGGC	TGCGGGCCTA	CTTTTTGGGC	ACCCACGACA
20701	CAGCGATTCC	CGGGCTTTCT	TGCCGGCGAC	AAGCTGGCCT	GCGCCATTGT	CAACACGGCC
20761	GGCCGCGAGA	CCGGAGGCGT	GCACTGGCTC	GCCTTCGGCT	GGAACCCGCG	CTCGCGCACC
20821	TGCTACATGT	TCGACCCCTT	TGGGTTCTCG	GACCGCCGGC	TCAAGCAGAT	TTACAGCTTC
20881	GAGTACGAGG	CCATGCTGCG	CCGAAGCGCC	GTGGCCTCTT	CGCCCGACCG	CTGTCTCAGC
		CCACCCAGAC				
21001	TGCATGTTCT	TGCATGCCTT	CGTGCACTGG	CCCGACCGAC	CCATGGACGG	GAACCCCACC
		TGACGGGGGT				
		ACCAGGAGGA				
		CCGCCATCGA				
		CAGCACTTTT				
		CTCGCGCTCA				
		CTGCCACTTG				
	-	CCACATACGC				
		GCAGTTGGGA				
		CACCATCAGA				
		GTCCAGATCC				
		GAATGGCACG				
		CGCGCCGCGC				
		TGTTGGGCCT				
		TTCCCGCACC				
		TGCCCGTTCT				
_		CCATGCAGAC				
		CCGGTGCACT				
		AGGCGACCCA				
		GCCTCCTCGT				
		AGCTTGTAAG				
		GTATCCATGC				
		AGGACACCGG				
		ACCGGAGGCT				
		TCGGGGTCTA				
		TCCACGGGA				
		TGGTGGGCAG				
		CCGACCCGTG				
		GCCGCCGGCC				
		GGAGGACTTA				
		GGCTCGTCTA				
		ACCGACGCTG				
		CTGCAGCGCC				
		AGCGTCGAGG				
		AAACGCCAGC				
		GCGGTCCCCG				
		TCCTGCCGCG				
		ATACCTGATA				
		AGACGCGCGC				
		CTGGTAGAGT				
		ACCCACTTCG				
23521	CATCATGGAT	CAGCTAATCA	TGCCCCACAT	CGAGGCCCTC	GATGAAAGTC	AGGAGCAGCG
23581	CCCCGAGGAC	ACCCGGCCCG	TGGTCAGCGA	TGAGCAGCTT	GCGCGCTGGC	TTGGTACCCG
		GCCCTGGAGC				
		TGCATGCGAC				
23761	ACCTGCACTA	CACTTTTAGC	ACGTTTCGTC	AGGCAGGCAT	GCAAGATCTC	CAACGTGGAG
23821	${\tt CTGACCAACT}$	GGTCTCCTGC	CTGGGAATCC	TGCACGAGAA	CCGCCTGGGG	CAGACAGTGC
23881	TCCACTCGAC	CCTGAAGGGC	GAGGCGCGGC	GGGACTATGT	CCGCGACTGC	GTCTTTCTCT

		CACATGGCAA				
24001	ACCTGAAGGA	GCTGGACAAG	CTTCTTGCTA	GAAACCTCAA	AAAGCTGTGG	ACGGGCTTTG
		CGTCGCCTCG				
		GCGGGCTGCC				
		AGCGATCTGG				
24241	GTCCCGCTGA	GCTACCGCGA	GTGTCCCCCG	CCGCTGTGGA	GCCACTGCTA	CCTCTTGCAG
24301	CTGGCCAACT	ACATCGCCTA	CCACTCGGAT	GTTATCGAGG	ACGTGAGCGG	CGAGGGGCTG
24361	CTAGAGTGCC	ACTGCCGCTG	CAACCTGTGC	TCTCCGCACC	GCTCCTGGTC	TGCAACCCCC
24421	AGCTCCTGAG	CGAGACCCAG	GTCATCGGTA	CCTTCGAGCT	GCAAGGTCCG	CAGGAGTCCA
24481	CCGCTCCGCT	GAAACTCACG	CCGGGGTTGT	GGACTTCCGC	GTACCTGCGC	AAATTTGTAC
24541	CCGAGGACTA	CCACGCCCAT	GAGATAAAGT	TCTTCGAGGA	CCAATCGCGC	CCGCAGCACG
24601	CGGATCTCAC	GGCCTGCGTC	ATCACCCAGG	GCGCGATCCT	CGCCCAATTG	CACGCCATCC
24661	AAAAATCCCG	CCAAGAGTTT	CTTTTGAAAA	AGGGTAGAGG	GGTCTATCTG	GACCCCCAGA
24721	CGGGCGAAGT	GCTCAACCCG	GGTCTCCCCC	AGCATGCCGA	AGAAGAACAG	GAGCCGCTAG
24781	TGGAAGAGAT	GGAAGAAGAA	TGGGACAGCC	AGCAGAAGAA	GACGAATGGG	AAGAAGAGAC
24841	AGAAGAAGAA	GAATTGGAAA	AGTGGAAGAA	GAGCAGCACA	${\tt GACACCGTCG}$	CCGCACCATC
24901	CGCGCCGCAG	CCCGGCGGTC	ACGGATACAA	CTCGCAGTCC	GCCAAGCTCC	TCGTAGATGG
24961	ATCGAGTGAA	GGTGACGGTA	AGCACGAGCG	${\tt GCAGGGCTAC}$	GAATCATGGA	GGCCCACAAA
25021	GCGGGATCAT	CGCCTGCTTG	CAAGACTGCG	GGGGGAACAT	${\tt CGTTTCGCCC}$	GCCGCTATCT
25081	GCTCTTCCAT	CGCGGGGTGA	ACATCCCCCG	CAACGTGTTG	CATTACTACC	${\tt GTCACCTTCA}$
	-	AAAATCAGAG				
25201	CGAGCCATTG	ACCACCAGGG	AGCTGAGGAA	TCGGATCTTC	CCCACTCTTT	ATGCCATTTT
25261	TCAGCAGAGT	CGAGGTCAGC	AGCAAGAGCT	CAAAGTAAAA	AACCGGTCTC	TGCGCTCGCT
25321	CACCCGCAGT	TGCTTGTACC	ACAAAAACGA	AGATCAGCTG	CAGCGCACTC	TCGAAGACGC
25381	CGAGGCTCTG	TTCCACAAGT	ACTGCGCGCT	CACTCTTAAA	GACTAAGGCG	CGCCCACCCG
25441	GAAAAAAGGC	GGGAATTACC	TCATCGCCAC	CATGAGCAAG	GAGATTCCCA	CCCCTTACAT
25501	GTGGAGCTAT	CAGCCCCAGA	TGGGCCTGGC	CGCGGGCGCC	TCCCAGGACT	ACTCCACCCG
25561	CATGAACTGG	CTCAGTGCCG	GCCCCTCGAT	GATCTCACGG	GTCAACGGGG	TCCGTAACCA
		ATATTGTTGG				
25681	CCGCGTAATT	GGCCCTCCAC	CCTGGTGTAT	CAGGAAATCC	CCGGGCCGAC	TACCGTACTA
25741	CTTCCGCGTG	ACGCACTGGC	CGAAGTCCGC	ATGACTAACT	CAGGTGTCCA	GCTGGCCGGC
25801	GGCGCTTCCC	GGTGCCCGCT	CCGCCCACAA	TCGGGTATAA	AAACCCTGGT	GATACGAGGC
25861	AGAGGCACAC	AGCTCAACGA	CGAGTTGGTG	AGCTCTTCAA	TCGGTCTGCG	ACCGGACGGA
		TAGCCGGAGC				
		CTCTTCGGAG				
26041	AGTTTGTGCC	CTCGGTCTAC	TTCAACCCCT	TCTCGGGATC	GCCAGGCCTC	TACCCGGACG
26101	AGTTCATACC	GAACTTCGAC	GCAGTGAGAG	AAGCGGTGGA	CGGCCACGAC	TGAATGTCTT
26161	ATGGTGACTC	GGCTGAGCTC	GCTCGGTTGA	GGCACCTAGA	CCACTGCCGC	CGCCTGCGCT*
		GGAGAGCTGC				
		CGGAGTGCGG				
26341	TCTTCACCCA	GCAACCCTTC	CTGGTCGAGC	GGGACCGGGG	AGGCACCACC	TACACCGTCT
26401	ACTGCATCTG	TCCAACCCCG	AAGTTGCATG	AGAATTTTTG	TTGTACTCTG	TGTGCTGAGT
		CTAAACTCCT				
		TCACCAACCA				
		AAAATTTTT				
		CCAACAACCT				
		ATCCTTTTGT				
		TGGTGAACGT				
26821	CTTTTTGATA	CTAACACTCC	TAAAACCGGA	GGTGAGCTCT	GGGTTCCCTC	TCTAACAGAG
		ATATTGAAGC				
26941	ATAGCGGTGC	TGTATTACCT	TCCTTGCTGG	ATCGAAATCA	AAATCTTTAT	CTGCTGGGTC
27001	AGACATTGTT	GGGAGGAACC	ATGAAGGGC	TCTTGCTGAT	TATCCTTTCC	CTGGTGGGGG
		ATGCCACGAA				
27121	TGATATGCAC	AGTAGTCATC	AAATGCGAGC	ATACATGCCC	TCTCAACATC	ACATTCAAAA
27181	ACCGTACCAT	GGGAAATGCA	TGGGTGGGCG	ACTGGGAACC	AGGAGATGAG	CAGAACTACA
27241	CGGTCACTGT	CCATGGTAGC	AATGGAAATC	ACACTTTTGG	TTTCAAATTC	ATTTTTGAAG
27301	TCATGTGTGA	TATCACACTG	CATGTGGCTA	GACTTCATGG	CTTGTGGCCC	CCTACCAAGG

27361 ATAACATGGT TGGGTTTTCT TTGGCTTTTG TGATCATGGC CTGTGCAATG TCAGGTCTGC 27421 TGGTAGGGC TTTAGTGTGG TTCCTAAAGC GCAAGCCTAG GTATGGAAAT GAGGAGAAGG 27481 AAAAATTGCT ATAAATCTTT TCTCTTCGCA GAACCATGAA TACAGTGATC CGTATCGTGC 27541 TGCTCTCT TCTTGTAACT TTTAGTCAGG CAGGATTCAT ACCATCAATG CTACATGGTG 27601 GGCTAATATA ACTTTAGTGG GACCTCAGAT ATTCCAGATC ACATGGTATG ATAGCACTGG 27661 ATTGCAATTT TGTGATGGAA GTACAGTTAA GAATCCACAG ATCAGACATA GTTGTAATGA 27721 TCAAAACTTA ACTCTGATTC ATGTGAACAA AACCCATGAA AGAACATACA TGGGCTATAA 27781 TAAGCAGAGT ACTCATAAAG AAGACTATAA AGTCACAGTT ATACCACCTC CTCCTGTTAC 27841 TGTAAAGCCA CAACCAGAGC CAGAATATGT GTATGTTAAT ATGGGAGAGA ACAAAACCTT 27901 AGTTGGGCCT CCAGGAATTC CAGTTAGTTG GTTTAATCAG GATGGTTTAC AATTTTGCAT 27961 TGGGGATAAA GTTTTCATC CAGAATTCAA CCACACCTGT GACATGCAAA ATCTTACACT 28021 GTIGTTTATA AATCTTACAC ATGATGGAGC TTATCTTGGT TATAATCGCC AGGGAACTGA 28081 AAGAACTTGG TATGAGGTTG TAGTGTCAGA TGGTTTTCCA AAATCAGAAG AGATGAAGGT 28141 AGAAGACCAT AGTAAAGAAA CAGAACAAAA ACAGACTGGT CAAAAACAAA GTGACCATAA 28201 GCAGGGTGGG CAAAAAGAAA CAAGTCAAAA GAAAACTAAT GACAAACAAA AGCCATCGCG 28261 CAGGAGGCCA TCTAAACTAA AGCCAAACAC ACCTGACACA AAACTAATTA CAGTCACTAG 28321 TGGGTCAAAC GTAACTTTAG TTGGTCCAGA TGGAAAGGTC ACTTGGTATG ATGATGATTT 28381 AAAAAGACCA TGTGAGCCTG GGTATAAGTT AGGGTGTAAG TGTGACAATC AAAACCTAAC 28441 CCTAATCAAT GTAACTAAAC TTTATGAGGG AGTTTACTAT GGTACTAATG ACAGAGGCAA 28501 CAGCAAAAGA TACAGAGTAA AAGTAAACAC TACTAATTCT CAAAGTGTGA AAATTCAGCC 28561 GTACACCAGG CCTACTACTC CTGATCAGAA ACACAGATTT GAATTGCAAA TTGATTCTAA 28621 TCAAGACAAA ATTCCATCAA CTACTGTGGC AATCGTGGTG GGAGTGATCG CGGGCTTTGT 28681 AACTCTAATC ATTATTTCA TATGCTACAT CTGCTGCCGC AAGCGTCCCA GGTCATACAA 28741 TCATATGGTA GACCCACTAC TCAGCTTCTC TTACTGAAAC TCAGTCACTC TCATTTCAGA 28801 ACCATGAAGG CTTTCACAGC TTGCGTTCTG ATTAGCATAG TCACACTTAG TTCAGCTGCA 28861 ATGATTAATG TTAATGTCAC TAGAGGTGGT AAAATTACAT TGAATGGGAC TTATCCACAA 28921 ACTACATGGA CAAGATATCA TAAAGATGGA TGGAAAAATA TTTGTGAATG GAATGTTACT 28981 GCATACAAAT GCTTCAATAA TGGAAGCATT ACTATTACTG CCACTGCCAA CATTACTTCT 29041 GGCACATACA AAGCTGAAAG CTATAAAAAT GAAATTAAAA AATTAACCTA TAAAAAACAAC 29101 AAAACCACAT TTGAAGATTC TGGAAATTAT GAGCATCAAA AATTATCTTT TTATATGTTG 29161 ACAATAATTG AACTGCCTAC AACCAAGGCA CCCACCACAG TTAGTACAAC TACACAGTCA 29221 ACTGTTAAGA CCACTACTCA CACTACACAG CTAGACACCA CAGTGCAGAA TAATACTGTG 29281 TTGGTTAGGT ATTTGTTGAG GGAGGAAAGT ACTACTGAAC AGACAGAGGC TACCTCAAGT 29341 GCCTTTATCA GCACTGCAAA TTTAACTTCG CTTGCTTGGA CTAATGAAAC CGGAGTATCA 29401 TTGATGCATG GCCAGCCTTA CTCAGGTTTG GATATTCAAA TTACTTTTCT GGTTGTCTGT 29461 GGGATCTTTA TTCTTGTGGT TCTTCTGTAC TTTGTCTGCT GTAAAGCCAG AAAGAAATCT 29521 AGGAGGCCCA TCTACAGGCC AGTGATTGGG GAACCTCAGC CACTCCAAGT GGATGGAGGC 29581 TTAAGGAATC TTCTTTCTC TTTTACAGTA TGGTGATCAG CCATGATTCC TAGTTCTTCC 29641 TATTTAACAT CCTCTTCTGT CTCTTCAACA TCTGTGCTGC CTTTGCGGCA GTTTCGCACG 29701 CCTCGCCCGA CTGTCTAGGG CCTTTCCCCA CCTACTCCTC TTTGCCCTGC TCACCTGCAC 29761 CTGCGTCTGC AGCATTGTCT GCCTGGTCAT CACCTTCCTG CAGCTCATCG ACTGGTGCTG 29821 CGCGCGCTAC AATTACTTCA TCATAGTCCC GAATACAGGG ACGAGAACGT AGCCAGAATT 29881 TTAAGGCTCA TATGACCATG CAGACTCTGC TCATACTGCT ATCGCTCTTA TCCCATGCCC 29941 TCGCTACTGC TGATTACTCT AAATGCAAAT TGGCGGACAT ATGGAATTTC TTAGACTGCT 30001 ATCAGGAGAA AATTGATATG CCCTCCTATT ACTTGGTGAT TGTGGGAATA GTTATGGTCT 30061 GCTCCTGCAC TTTCTTTGCC ATCATGATCT ACCCCTGTTT TGATCTTGGA TGGAACTCTG 30121 TTGAGGCATT CACATACACA CTAGAAAGCA GTTCACTAGC CTCCACGCCA CCACCCACAC 30181 CGCCTCCCCG CAGAAATCAG TTTCCCATGA TTCAGTACTT AGAAGAGCCC CCTCCCCGAC 30241 CCCCTTCCAC TGTTAGCTAC TTTCACATAA CCGGCGGCGA TGACTGACCA CCACCTGGAC 30301 CTCGAGATGG ACGGCCAGGC CTCCGAGCAG CGCATCCTGC AACTGCGCGT CCGTCAGCAG 30361 CAGGAGCGTG CCGCCAAGGA GCTCCTCGAT GCCATCAACA TCCACCAGTG CAAGAAGGGC 30421 ATCTTCTGCC TGGTCAAACA GGCAAAGATC ACCTACGAGC TCGTGTCCAA CGGCAAACAG 30481 CATCGCCTCA CCTATGAGAT GCCCCAGCAG AAGCAGAAGT TCACCTGCAT GGTGGGCGTC 30541 AACCCCATAG TCATCACCCA GCAGTCGGGC GAGACCAACG GCTGCATCCA CTGCTCCTGC 30601 GAAAGCCCCG AGTGTATCTA CTCCCTTCTC AAGACCCTTT GCGGACTCCG CGACCTCCTC 30661 CCCATGAACT GATGTTGATT AAAAACCAAA AAAAACAATC AGCCCCTTCC CCTATCCCAA 30721 ATTACTCGCA AAAATAAATC ATTGGAACTA ATCATTTAAT AAAGATCACT TACTTGAAAT

30781 CTGAAAGTAT GTCTCTGGTG TAGTTGTTCA GCAGCACCTC GGTACCCTCC TCCCAACTCT 30841 GGTACTCCAG TCTCCGGCGG GCGGCGAACT TTCTCCACAC CTTGAAAGGG ATGTCAAATT 30901 CCTGGTCCAC AATTTCATT GTCTTCCCTC TCAGATGTCA AAGAGGCTCC GGGTGGAAGA 30961 TGACTTCAAC CCCGTCTACC CCTATGGCTA CGCGCGGAAT CAGAATATCC CCTTCCTCAC 31021 TCCCCCCTTT GTCTCCTCCG ATGGATTCAA AAACTTCCCC CCTGGGGTCC TGTCACTCAA 31081 ACTGGCTGAC CCAATCACCA TAGCCAATGG TGATGTCTCA CTCAAGGTGG GAGGGGACTT 31141 ACTTTGCAAG AAGGAAGTAT GACTGTAGAC CCTAAGGCTC CCTTGCAACT TGCAAACAAT 31201 AAAAAACTTG AGCTTGTTTA TGTTGATCCA TTTGAGGTTA GTGCCAATAA ACTTAGTTTA 31261 AAAGTAGGAC ATGGATTAAA AATATTAGAT GACAAAAGTG CTGGAGGGTT GAAAGATTTA 31321 ATTGGCAAAC TTGTGGTTTT AACAGGGGAA AGGAATAGGC ACTGAAAATT TGCAAAATAC 31381 AGATGGTAGC AGCAGAGGAA TTGGTATAAG TGTAAGAGCA AGAGAAGGGT TAACATTTGA 31441 CAATGATGGA TACTTGGTAG CATGGAACCC AAAGTATGAC ACGCGCACAC TTTGGACAAC 31501 ACCAGACACA TCTCCTAATT GCAGGATTGA TAAGGAGAAG ATTCAAAACT CACTTTGGTA 31561 CTTACAAAGT GTGGAAGTCA AATATTAGCT AATGTGTCTT TGATTGTGGT GTCAGGAAAA 31621 TATCAATACA TAGACCACGC TACAAATCCA ACTCTTAAAAT CATTTAAAAT AAAACTTCTT 31681 TTTGATAATA AAGGTGTACT TCTCCCAAGT TCAAACCTTG ATTCCACATA TTGGAACTTT 31741 AGAAGTGACA ATTTAACTGT ATCTGAGGCA TATAAAAATG CAGTTGAATT TATGCCTAAT 31801 TTGGTAGCCT ACCCAAAACC TACCACTGGC TCTAAAAAAT ATGCAAGGGA TATAGTCTAT 31861 GGGAACATAT ATCTTGGAGG TTTGGCATAT CAGCCAGTTG TAATTAAGGT TACTTTTAAT 31921 GAAGAAGCAG ATAGTGCTTA CTCTATAACA TTTGAATTTG TATGGAATAA AGAATATGCC 31981 AGGGTTGAAT TTGAAACCAC TTCCTTTACC TTCTCCTATA TTGCCCAACA ATAAAAGACC 32041 AATAAACGTG TTTTTTATTT CAAATTTTAT GTATCTTTAT TGATCTTTAC ACCAGCGCGA 32101 GTAGTCAATC TCCCACCACC AGCCCATTTC ACAGTGTACA CGGTTCTCTC AGCACGGTGG 32161 CCTTAAATAA GGAAATGTTC TGATTATTGC GGGAACTGGA CTTGGGGTCT ATAATCCACA 32221 CAGTTTCCTG ACGAGCCAAA CGGGGATCGG TGATTGAAAT GAAGCCGTCC TCTGAAAAGT 32281 CATCCAAGCG GGCCTCACAG TCCAGGTCAC AGTCTGGTGG AACGAGAAGA ACGCACAGAT 32341 TCATACTCGG AAAACAGGAT GGGTCTGTGC CTCTCCATCA GCGCCCTCAG CAGTCTCTGC 32401 CGCCGGGGCT CGGTGCGGCT GCTGCAAATG GGATCGGGAT CACAAGTCTC TCTAACTATG 32461 ATCCCAACAG CCTTCAGCAT CAGTCTCCTG GTGCGTCGAG CACAGCACCG CATCCTGATC 32521 TCTGCCATGT TCTCACAGTA AGTGCAGCAC ATAATCACCA TGTTATTCAG CAGCCCATAA 32581 TTCAGGGTGC TCCAGCCAAA GCTCATGTTG GGGATGATGG AACCCACGTG ACCATCGTAC 32641 CAGATGCGGC AGTATATCAG GTGCCTGCCC CTCATGAACA CACTGCCCAT ATACATGATC 32701 TCTTTGGGCA TGTTTCTGTT TACAATCTGG CGGTACCAGG GGAAGCGCTG GTTGAACATG 32761 CACCCGTAAA TGACTCTCCT GAACCACACG GCCAGCAGGG TGCCTCCCGC CCGACACTGC 32821 AGGGAGCCAG GGGATGAACA GTGGCAATGC AGGATCCAGC GCTCGTACCC GCTCACCATC 32881 TGAGCTCTTA CCAAGTCCAG GGTAGCGGGG CACAGGCACA CTGACATACA TCTTTTTAAA 32941 ATTTTATTT CCTCTGTGGT GAGGATCATA TCCCAGGGGA CTGGAAACTC TTGGAGCAGG 33001 GTAAAGCCAG CAGCACATGG TAATCCACGG ACAGAACTTA CATTATGATA ATCTGCATGA 33061 TCACAATCGG GCAACAGGGG ATGTTGATCA GTCAGTGAAG CCCTGGTTTC ATCATCAGAT 33121 CGTGGTAAAC GGGCCCTGCG ATATGGATGA TGGCGGAGCG AGCTGGATTG AATCTCGGTT 33181 TGCATTGTAG TGGATTCTCT TGCGTACCTT GTCGTACTTC TGCCAGCAGA AATGGGCCCT 33241 TGAACAGCAT ATACCCCTCC TGCGGCCGTC CTTTCGCTGC TGCCGCTCAG TCATCCAACT 33301 GAAGTACATC CATTCTCGAA GATTCTGGAG AAGTTCCTCT GCATCTGATG AAATAAAAAA 33361 CCCGTCCATG CGAATTCCCC TCATCACATC AGCCAGGACT CTGTAGGCCA TCCCCATCCA 33421 GTTAATGCTG CCTTGTCTAT CATTCAGAGG GGGCGGTGGC AGGATTGGAA GAACCATTTT 33481 TATTCCAAAC GGTCTCGAAG GACGATAAAG TGCAAGTCAC GCAGGTGACA GCGTTCCCCT 33541 CCGCTGTGCT GGTGGAAACA GACAGCCAGG TCAAAACCCA CTCTATTTTC AAGGTGCTCG 33601 ACCGTGGCTT CGAGCAGTGG CTCTACGCGT ACATCCAGCA TAAGAATCAC ATTAAAGGCT 33661 GGCCCTCCAT CGATTTCATC AATCATCAGG TTACATTCCT GCACCATCCC CAGGTAATTC 33721 TCATTTTCC AGCCTTGGAT TATCTCTACA AATTGTTGGT GTAAATCCAC TCCGCACATG 33781 TTGAAAAGCT CCCACAGTGC CCCCTCCACT TTCATAATCA GGCAGACCTT CATAATAGAA 33841 ACAGATCCTG CTGCTCCACC ACCTGCAGCG TGTTCAAAAC AACAAGATTC AATAAGGTTC 33901 TGCCTCCGC CCTGAGCTCG CGCCTCAATG TCAGCTGCAA AAAGTCACTT AAGTCCTGGG 33961 CCACTACAGC TGACAATTCA GAGCCAGGGC TAAGCGTGGG ACTGGCAAGC GTGAGGGAAA 34021 ACTTTAATGC TCCAAAGCTA GCACCCAAAA ACTGCATGCT GGAATAAGCT CTCTTTGTGT 34081 CTCCGGTGAT GCCTTCCAAA ATGTGAGTGA TAAAGCGTGG TAGTTTTTTC TTTAATCATT 34141 TGCGTAATAG AAAAGTCCTG TAAATAAGTC ACTAGGACCC CAGGGACCAC AATGTGGTAG

34201	CTTACACCGC	GTCGCTGAAA	GCATGGTTAG	TAGAGATGAG	AGTCTGAAAA	ACAGAAAGCA
34261	TGCGCTAAAC	TAAGGTGGCT	ATTTTCACTG	AAGGAAAAAT	CACTCTTTCC	AGCAGCAGGG
34321	TACCCACTGG	GTGGCCCTTG	CGGACATACA	AAAATCGGTC	CGTGTGATTA	AAAAGCAGCA
34381	CAGTAAGTTC	CTGTCTTCTT	CCGGCAAAAA	TCACATCGGA	CTGGGTTAGT	ATGTCCCTGG
34441	CATGGTAGTC	ATTCAAGGCC	ATAAATCTGC	CCTGATATCC	AGTAGGAACC	AGCACACTCA
34501	CTTTTAGGTG	AAGCAATACC	ACCCCATGCG	${\tt GAGGAATGTG}$	GAAAGATTCA	GGGCAAAAA
34561	AATTATATCT	ATTGCTAGCC	CTTCCTGGAC	${\tt GGGAGCAATC}$	CTCCAGGACT	ATCTATGAAA
34621	GCATACAGAG	ATTCAGCCAT	AGCTCAGCCC	GCTTACCAGT	AGACAAAGAG	CACAGCAGTA
34681	CAAGCGCCAA	CAGCAGCGAC	TGACTACCCA	CTGACTTAGC	TCCCTATTTA	AAGGCACCTT
34741	ACACTGACGT	AATGACCAAA	${\tt GGTCTAAAAA}$	CCCCGCCAAA	AAAACACACA	CGCCCTGGGT
34801	GTTTTTGCGA	AAACACTTCC	GCGTTCTCAC	TTCCTCGTAT	CGATTTCGTG	ACTTGACTTC
34861	CGGGTTCCCA	CGTTACGTCA	${\tt CTTTTGCCCT}$	TACATGTAAC	TTAGTCGTAG	GGCGCCATCT
34921	TGCCCACGTC	${\tt CAAAATGGCT}$	TACATGTCCA	GTTACGCCTC	CGCGGCGACC	GTTAGCCGTG
34981	CGTCGTGACG	TCATTTGCAT	CAACGTTTCT	CGGCCAATCA	GCAGTAGCCC	CGCCCTAAAT
35041	TTAAAACCTC	ATTTGCATAT	TAACTTTTGT	TTACTTTGTG	GGGTATATTA	TTGATGATG

ATGTCAAAGAGGCTCCGGGTGGAAGATGACTTCAACCCCGTCTACCCCTA TGGCTACGCGCGAATCAGAATATCCCCTTCCTCACTCCCCCCTTTGTCTC CTCCGATGGATTCAAAAACTTCCCCCCTGGGGTCCTGTCACTCAAACTGGC TGACCCAATCACCATAGCCAATGGTGATGTCTCACTCAAGGTGGGAGGGG GACTTACTTTGCAAGAAGGAAGTCTGACTGTAGACCCTAAGGCTCCCTTG CAACTTGCAAACAATAAAAAACTTGAGCTTGTTTATGTTGATCCATTTGAG GTTAGTGCCAATAAACTTAGTTTAAAAGTAGGACATGGATTAAAAATATT AGATGACAAAAGTGCTGGAGGGTTGAAAGATTTAATTGGCAAACTTGTGG TTTTAACAGGGAAAGGAATAGGCACTGAAAATTTGCAAAATACAGATGGT AGCAGCAGAGGAATTGGTATAAGTGTAAGAGCAAGAGAAAGGGTTAACAT TTGACAATGATGGATACTTGGTAGCATGGAACCCAAAGTATGACACGCGC ACACTTTGGACAACACCAGACACATCTCCTAATTGCAGGATTGATAAGGA GAAGGATTCAAAACTCACTTTGGTACTTACAAAGTGTGGAAGTCAAATAT TAGCTAATGTGTCTTTGATTGTGGTGTCAGGAAAATATCAATACATAGACC ATAAAGGTGTACTTCTCCCAAGTTCAAACCTTGATTCCACATATTGGAACT TTAGAAGTGACAATTTAACTGTATCTGAGGCATATAAAAATGCAGTTGAA TTTATGCCTAATTTGGTAGCCTACCCAAAACCTACCACTGGCTCTAAAAAA TATGCAAGGGATATAGTCTATGGGAACATATATCTTGGAGGTTTGGCATA TCAGCCAGTTGTAATTAAGGTTACTTTTAATGAAGAAGCAGATAGTGCTTA CTCTATAACATTTGAATTTGTATGGAATAAAGAATATGCCAGGGGTTGAA TTTGAAACCACTTCCTTTACCTTCTCCTATATTGCCCAACAATAA

SEQ ID NO:2

Penton17.Seq Length: 1554

1 ATGAGGCGTG CGGTGGTGTC TTCCTCTCCT CCTCCCTCGT ACGAGAGCGT 51 GATGGCGCAG GCGACCCTGG AGGTTCCGTT TGTGCCTCCG CGGTATATGG CTCCTACGGA GGGCAGAAAC AGCATTCGTT ACTCGGAGCT GGCTCCGTTG 101 TACGACACCA CTCGCGTGTA CTTGGTGGAC AACAAGTCGG CGGACATCGC 151 TTCCCTGAAC TATCAAAACG ACCACAGCAA CTTCCTGACC ACGGTGGTGC 201 251 AGAACAACGA TTTCACCCCC GCCGAGGCTA GCACGCAGAC GATAAATTTT 301 GACGAGCGGT CGCGGTGGGG CGGTGATCTG AAGACCATTC TGCACACCAA 351 CATGCCCAAT GTGAACGAGT ACATGTTCAC CAGCAAGTTT AAGGCGCGGG TGATGGTGGC TAGAAAACAC CCACAGGGGG TAGAAGCAAC AGATTTAAGC 401 451 AAGGATATCT TAGAGTATGA GTGGTTTGAG TTTACCCTGC CCGAGGGCAA 501 CTTTTCCGAG ACCATGACCA TAGACCTGAT GAACAACGCC ATCTTGGAAA 551 ACTACTTGCA AGTGGGGCGG CAAAATGGCG TGCTGGAGAG CGATATTGGA 601 GTCAAGTTTG ACAGCAGAAA TTTCAAGCTG GGCTGGGACC CTGTGACCAA 651 GCTGGTGATG CCAGGGGTCT ACACCTACGA GGCCTTTCAC CCGGACGTGG 701 TGCTGCTGCC GGGCTGCGGG GTGGACTTCA CAGAGAGCCG CCTGAGCAAC CTCCTGGGCA TTCGCAAGAA GCAACCTTTC CAAGAGGGCT TCAGAATCAT 751 GTATGAGGAT CTAGAAGGGG GCAACATCCC CGCCCTGCTG GATGTGCCCA 801 851 AGTACTTGGA AAGCAAGAAG AAGTTAGAGG AGGCATTGGA GAATGCTGCT 901 AAAGCTAATG GTCCTGCAAG AGGAGACAGT AGCGTCTCAA GAGAGGTTGA AAAGGCAGCT GAAAAAGAAC TTGTTATTGA GCCCATCAAG CAAGATGATA 951 1001 CCAAGAGAAG TTACAACCTC ATCGAGGGAA CCATGGACAC GCTGTACCGC 1051 AGCTGGTACC TGTCCTATAC CTACCGGGAC CCTGAGAACG GGGTGCAGTC 1101 GTGGACGCTG CTCACCACCC CGGACGTCAC CTGCGGCGCG GAGCAAGTCT 1151 ACTGGTCGCT GCCGGACCTC ATGCAAGACC CCGTCACCTT CCGTTCTACC 1201 CAGCAAGTCA GCAACTACCC CGTGGTCGGC GCCGAGCTCA TGCCCTTCCG 1251 CGCCAAGAGC TTTTACAACG ACCTCGCCGT CTACTCCCAG CTCATCCGCA 1301 GCTACACCTC CCTCACCCAC GTCTTCAACC GCTTCCCCGA CAACCAGATC

SEQ ID NO: 3

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1351	CTCTGCCGTC	CGCCCGCGCC	CACCATCACC	ACCGTCAGTG	AAAACGTGC
L 401 .	TGCTCTCACA	GATCACGGGA	CGCTACCGCT	GCGCAGCAGT	ATCCGCGGA
L 451	TCCAGCGAGT	GACCGTCACT	GACGCCCGTC	GCCGCACCTG	TCCCTACGT
501	TACAAGGCCC	TGGGCATAGT	CGCGCCGCGT	GTGCTTTCCA	GTCGCACCT"
1551	CTAA				

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Claims

- A chimeric adenoviral vector comprising nucleotide sequence of a first adenovirus, wherein at least one gene of said first adenovirus encoding a protein that facilitates binding of said vector to a target mammalian cell, or internalization thereof within said cell, is replaced by the corresponding gene from a second adenovirus belonging to subgroup D, said vector further comprising a transgene operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell.
- A chimeric adenoviral vector according to Claim 1 wherein said second adenovirus is selected from the group consisting of Ad 9, Ad 15, Ad 17, Ad 19, Ad 20, Ad 22, Ad 26, Ad 27, Ad 28, Ad 30, and Ad 39.
- 15 3. A chimeric adenoviral vector according to Claim 1 wherein said first adenovirus is selected from the group consisting of Ad 2, Ad 5, and Ad 12.
 - 4. A chimeric adenoviral vector according to Claim 1 wherein said replaced gene encodes Ad fiber.
 - A chimeric adenoviral vector according to Claim 1 wherein said replaced gene encodes Ad penton base.
- 6. A chimeric adenoviral vector according to Claim 1 wherein a first replaced gene encodes Ad fiber, and a second replaced gene encodes Ad penton base.
 - 7. A chimeric adenoviral vector comprising nucleotide sequence of a first adenovirus, wherein a portion of a gene thereof encoding a protein that facilitates binding of said vector to a target mammalian cell, or internalization

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thereof within said cell, is replaced by a portion of the corresponding gene from a second adenovirus belonging to subgroup D, said vector further comprising a transgene operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell.

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- A chimeric adenoviral vector according to Claim 7 wherein the encoding 8. sequence that is replaced codes for a portion of Ad fiber.
- A chimeric adenoviral vector according to Claim 7 wherein the encoding 9. sequence that is replaced codes for a portion of Ad penton base. 10
 - A chimeric adenoviral vector according to Claim 9 wherein the encoding 10. sequence that is replaced codes for an amino acid sequence that includes RGD.
- A method of providing a biologically active protein to the airway epithelial 15 11. cells of a patient comprising administering to said cells an adenoviral vector selected from the group consisting of:

20

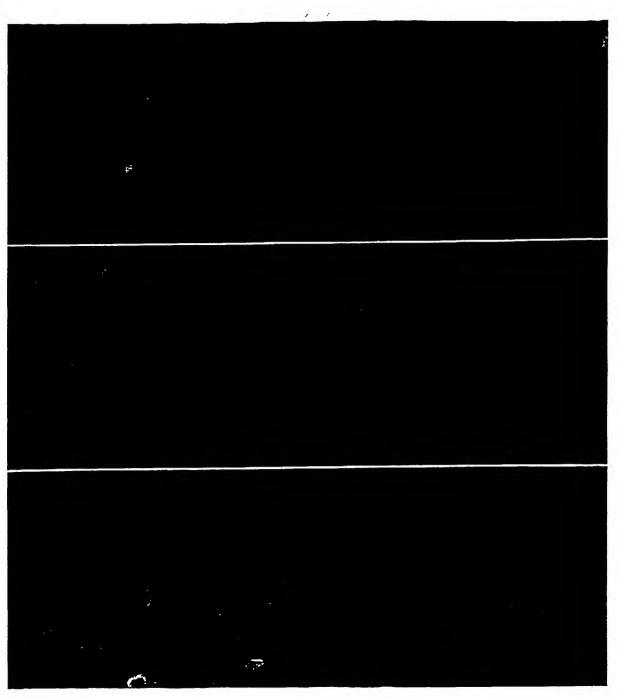
25

- (a) a chimeric adenoviral vector comprising nucleotide sequence of a first adenovirus, wherein at least one gene of said first adenovirus encodes a protein that facilitates binding of said vector to a target mammalian cell, or internalization thereof within said cell, is replaced by the corresponding gene from a second adenovirus belonging to subgroup D, said vector further comprising a transgene encoding said protein that is operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell; and
- (b) a chimeric adenoviral vector comprising nucleotide sequence of a first adenovirus, wherein a portion of a gene thereof encoding a protein that facilitates binding of said vector to a target mammalian cell, or internalization thereof within said cell, is replaced by a portion of the

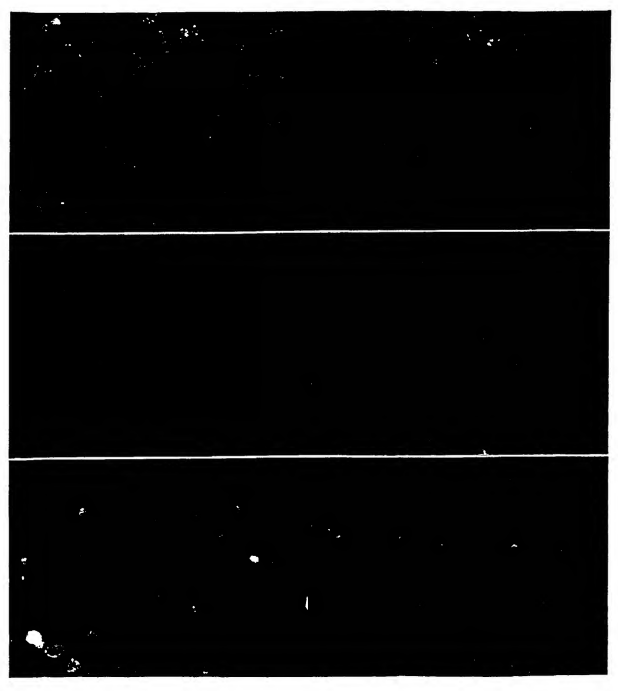
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corresponding gene from a second adenovirus belonging to subgroup D, said vector further comprising a transgene encoding said protein that is operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell;

- under conditions whereby the transgene encoding said protein is expressed, and phenotypic benefit is produced in said airway epithelial cells.
- 12. A method according to Claim 11 wherein said second adenovirus is Ad 17 and the nucleotide sequence thereof used in replacement of nucleotide sequence of said first adenovirus encodes a polypeptide selected from the group consisting of Ad 17 fiber, a fragment of Ad 17 fiber, Ad 17 hexon, a fragment of Ad 17 hexon, Ad penton base, and a fragment of Ad 17 penton base.
- 13. A method of providing a biologically active protein to the airway epithelial cells of a patient that comprises administering to said cells an adenoviral vector comprising elements of an Ad 17 genome, and a transgene encoding said protein that is operably linked to a eucaryotic promoter to allow for expression therefrom in a mammalian cell, under conditions whereby the transgene encoding said protein is expressed, and phenotypic benefit is produced in said airway epithelial cells.



F16 1 - original filed in PTO 15 Sull color - see side Solder



F162 - original 5. Cod in FTO
15 Sull colon - see 5.1 de 50 lder

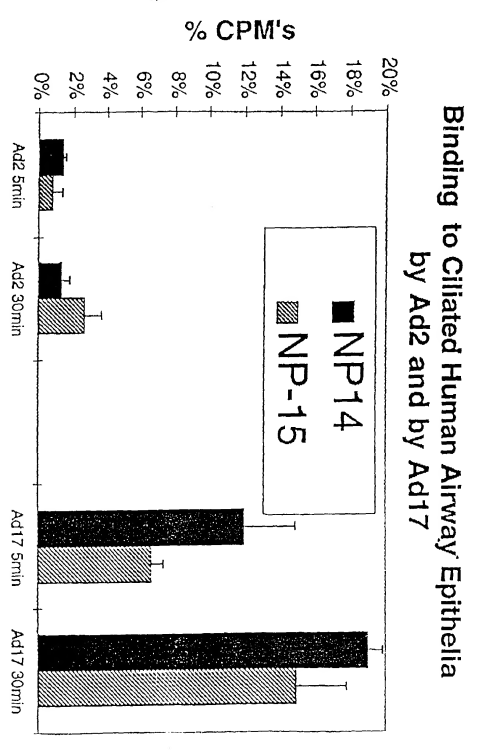
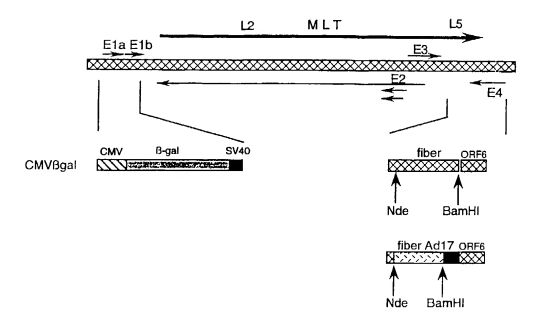


FIGURE 3

Chimeric Ad2/ßgal-2/ Ad17 vectors



M.RRAVVSSSPPPSYESVMAQATLEVPFVPRYMAPTEGR 39 : : : . : MQ.RAAMYEEGPPPSYESVVSAAPVAAALGSPFDAPLDPPFVPRYLRPTGGR 52	SEA ID NO:4 SEA ID NO:5
40 NSIRYSELAPLYDTTRVYLVDNKSADIASLNYQNDHSNFLTTVVONNDFT 89	
90 PAEASTOTINFDERSRWGGDLKTILHTNMPNVNEYMFTSKFKARVMVARK 139	
140 HPQGVEATDLSKDILEYEWFEFTLPEGNFSETMTIDLMNNAILENYLQVG 189	
190 RONGVLESDIGVKFDSRNFKLGWDPVTKLVMPGVYTYEAFHPDVVLLPGC 239	
240 GVDFTESRLSNLLGIRKKOPFOEGFRIMYEDLEGGNIPALLDVPKYLES. 288	— START
289 KKKLEEALENAAKANGPA	
314 REVEKAAE	
344 TMD.TLYRSWYLSYTYRDPENGVQSWTLLTTPDVTCGAEQVYWSLPDLMQ 392	
1	
END	

393	DPVTFRSTQQVSNYPVVGAELMPFRAKSFYNDLAVYSQLIRSYTSLTHVF	442
447	DPVTFRSTSQISNFPVVGAELLPVHSKSFYNDQAVYSQLIRQFTSLTHVF	496
443	NRFPDNQILCRPPAPTITTVSENVPALTDHGTLPLRSSIRGVQRVTVTDA	492
497	NRFPENQILARPPAPTITTVSENVPALTDHGTLPLRNSIGGVQRVTITDA	54 6
493	RRRTCPYVYKALGIVAPRVLSSRTF 517	
547	RRRTCPYVYKALGIVSPRVLSSRTF 571	

FIGURE SB

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//
         ...mrraam. ....yeegp ppsyesvvsa ..apvaaalg spfdapldpp 👉 SEQ ID NO: 6
 Penton5
          ...MORAAM. ....YEEGP PPSYESVVSA ..APVAAALG SPFDAPLDPP +5EQ ID NO:5
  Penton2
          ...MRRRAVLG GAV.VYPEGP PPSYESVM......QQQA AMIQPPLEAP - SEA ID NO: 7
 Penton3
Penton12 ...MRRAVEL QTV.AFPETP PPSYETVM. .....AAAPP 560 ID NO: 8
Penton40 ...MRRAVGV PPVMAYAEGP PPSYESVM.....ET ADLPATIQAL SEG ID NO: 9
         Penton17
Penton17 ...MRRAVV. .....SSSF FFSTESVIII. .....
Pentonf10 MWGLQPPTSI PPPPPPTELT PSTYPAMVNG YPPPAASAQS CSSSGGOSEL SGQ ID NO:10
 Penton5 FVP.PRYLRP TGGRNSIRYS ELAPLFDTTR VYLVDNKSTD VASLNYQNDH
          FVP.PRYLRP TGGRNSIRYS ELAPLFDTTR VYLVDNKSTD VASLNYQNDH
 Penton3 FVP.PRYLAP TEGRNSIRYS DVSPLYDTTK LYLVDNKSAD IASLNYQNDH
Penton12 YVP.PRYLGP TEGRNSIRYS ELSPLYDTTR VYLVDNKSSD IASLNYQNDH
Penton40 HVP.PRYLGP TEGRNSIRYS ELAPLYDTTR VYLVDNKSAD IASLNYQNDH
          FVP. PRYMAP TEGRNSIRYS ELAPLYDTTR VYLVDNKSAD IASLNYONDH
Penton17
Pentonf10 YMPLQRVMAP TGGRNSIKYR DYTPCRNTTK LFYVDNKASD IDTYNKDANH
          101
          SNFLTTVION NDYSPGEAST QTINLDDRSH WGGDLKTILH TNMPNVNEFM
 Penton5
          SNFLTTVIQN NDYSPGEAST QTINLDDRSH WGGDLKTILH TNMPNVNEFM
 Penton2
         SNFLTTVVON NDFTPTEAST OTINFDERSR WGGOLKTIMH TNMPNVNEYM
 Penton3
Penton12 SNFLTTVVQN NDYSPIEAGT QTINFDERSR WGGDLKTILH TNMPNVNDFM
Penton40 SNFQTTVVQN NDFTPTEAGT QTINFDDRSR WGGDLKTILR TNMPNINEFM
Penton17
          SNFLTTVVQN NDFTPAEAST QTINFDERSR WGGDLKTILH TNMPNVNEYM
Pentonf10 SNFRTTVIHN QDLDADTAAT ESIQLDNRSC WGGDLKTAVR TNCPNVSSFF
          151
 Penton5
          FTNKFKARVM VSRL..... PTKD..N QVELKYEWVE FTLPEGNYSE
 Penton2 FTNKFKARVM VSRS..... ...LTKD..K QVELKYEWVE FTLPEGNYSE
 Penton3 FSNKFKARVM VSRKAPEGVT VNDTYDH..K EDILKYEWFE FILPEGNFSA
Penton12 FTTKFKARVM VARK..... TNNE..G QTILEYEWAE FVLPEGNYSE
Penton40 STNKFRARVM VEK......VNR.K TNAPRYEWFE FTLPEGNYSE
Penton17 FTSKFKARVM VARKHPQGV. ..EATDL.S KDILEYEWFE FTLPEGNFSE
Pentonf10 QSNSVRVRMM WKRDPPTSTA PPSAVGSGYS VPGAQYKWYD LTVPEGNYAL
 Penton5 TMTIDLMNNA IVEHYLKVGR ONGVLESDIG VKFDTRNFRL GFDPVTGLVM
```

FIGURE GA

Penton2	TMTIDLMNNA	IVEHYLA-GR	ONGVLESDIC	VKFDTRNFRL	GFDPVIY
Penton3	TMTIDLMNNA	IIDNYLEIGR	ONGVLESDIC	VKFDTRNFRL	GWDPETKI.TM
Penton12	TMTIDLMNNA	IIEHYLRVGR	OHGVLESDIG	VKFDTRNFRL	GWDPETOLUT
Penton40	TMTIDLMNNA	IVDNYLAVGR	ONGVLESDIG	VKFDTRNFRL	GWDPVTKLVM
Penton17	TMTIDLMNNA	ILENYLOVGR	ONGVLESDIG	VKFDSRNFKL	GWDPVTKI.VM
Pentonf10	CELIDLLNEG	IVOLYLSEGR	ONNVOKEDTO	VKFDTRNFGL	I.RDPWWittwm
		1182120201	Z-MAZMODIC	VIG DITMINGE	INDEA1GEAT
	251				300
Penton5	PGVYTNEAFH	PDIILLPGCG	VDFTHSRLSN	LLGIRKROPF	
Penton2	PGVYTNEAFH	PDIILLPGCG	VDFTHSRLSN	LLGIRKROPF	OFCERTAND
Penton3	PGVYTYEAFH	PDIVLLPGCG	VDFTESRLSN	LLGIRKRHPF	OFCEKTAVED
Penton12	PGVYTNEAFH	PDTVLLPGCG	VDFTESRI.SN	ILGIRKROPF	OFCENTAMEN
Penton40	PGVYTNEAFH	PDTVLLPGCG	VDFTOSRI NN	LLGIRKRMPF	OCCEOTMORE
Penton17	PGVYTYEAFH	PDVVI.I.PGCG	UDFTESPI.SN	LLGIRKKOPF	ORGEDIAMEN
Pentonf10	PGTYVYKGYH	POTVILLEGEA	Thervenier	LLGIGKREPY	CVCENTER
1 checonita	10111111111	IDIVIDICA	IDF I I SKLISL	LIGIGRREPI	SKGFVITTED
	301				350
Penton5	LEGGNIPALL	DVDAYOASLK	DDTEOGGGGA	GGSNSSGSGA	
Penton2	LEGGNIPALL	DVDAYOASLK	DDTEOGGDGA	GGGNNSGSGA	EENSNAAAA
Penton3	LEGGNIPALL	DVTAYEESKK	A. זיזיינייניינייניינייניינייניינייניינייני	VAEETSE	
Penton12	LEGGNIPALL	DVKKYENSI.		Q	
Penton40	LEGGNIPALL	DVEKYEASTK			
Penton17	LECCHIPALI.	MORALECKA	KIE E	ALENAAK	
Pentonf10	LOGGDTPALL	DIDSTITUTE	DEPUTE DAY	A	
	DQOODIIADD	DIDDOVDVIIDA	DOEVIEDDIA	A	•••••
	351				400
Penton5	MOPVEDMNDH	AIRGDTFATR	AEEKRAEAEA	AAEAAAPAAQ	
Penton2	MOPVEDMNDH	AIRGDTFATR	AEEKRAEAEA	AAEAAAPAAQ	PEVEKPOKKP
Penton3	DDD	ITRGDTYITE	KOKREAAAAE	v	KKEI.
Penton12	DON	TVRGDNFIA.		L,	NKAA
Penton40		ETRGADEKEN	PΩ		חחווווווווווווווווווווווווווווווווווווו
Penton17	ANG	PARGDSSVSR	EVEKAA		EVET
Pentonf10			DVDIOUT		·····EREL
				• • • • • • • • • • • • • • • • • • • •	
	401				450
Penton5	VIKPLTEDSK	KRSYNLI	SNDSTFTQYR	SWYLAYNYGD	POTGIRSWTL
Penton2	VIKPLTEDSK	KRSYNLI	SNDSTFTQYR	SWYLAYNYGD	POTGIRSWTL
Penton3	KIQPLEKDSK	SRSYNVL	E.DKINTAYR	SWYLSYNYGN	PEKGIRSWTL
Penton12				SWYLAYNYGD	
Penton40				SWFLAYNYGD	
Penton17	VIEPIKQDDT	KRSYNLI	E.GTMDTLYR	SWYLSYTYRD	PENGVOSWTL
Pentonf10	PLLHDSA	GVSYNVIYDQ	VTGKPVTAYR	SWMLAYNVPN	SOANOTTL
	451				500
Penton5	LCTPDVTCGS	EQVYWSLPDM	MODPVTFRST	RQISNFPVVG	AELLPVHSKS
Penton2	LCTPDVTCGS	EQVYWSLPDM	MODPVTFRST	SQISNFPVVG	AELLPVHSKS
Penton3	LTTSDVTCGA	EOVYWSLPDM	MODPVTFRST	ROVNNYPVVG	AELMPVFSKS
Penton12	LTTPDVTGGS	EQVYWSLPDM	MODPVTFRSS	ROVSNYPVVA	AELLPVHAKS
Penton40				TOVSNYPVVG	
Penton17	LTTPDVTCGA	EQVYWSLPDL	MODPVTFRST	QQVSNYPVVG	AELMPFRAKS
Pentonf10	LTVPDMAGGI	GAMYTSLPDT	FIAPTGFKED	NTTNLCPVVG	MNLFPTYNKI
Dort- 5	501				550
Penton5	PYNDQAVYSQ	LIROFT.SLT	HVFNRFPENQ	ILARPPAPTI	TTVSENVPAL
Penton2	FYNDQAVYSQ	LIRQFT.SLT	HVFNRFPENQ	ILARPPAPTI	TTVSENVPAL
Penton3	FYNEQAVYSQ	QLRQAT.SLT	HVFNRFPENQ	ILIRPPAPTI	TTVSENVPAL
Penton12	FYNEQAVYSQ	LIRQST.ALT	RVFNRFPENQ	ILVRPPAATI	TTVSENVPAL

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Penton40	FYNEQAVYSQ	LIROST.ALT	HIFNRFPENQ	ILVRPPAPTI	TTVSENVFAL
Penton17	FYNDLAVYSQ	LIRSYT.SLT	HVFNRFPDNQ	ILCRPPAPTI	TTVSENVPAL
Pentonf10	YYQAASTYVQ	RLENSCQSAT	AAFNRFPENE	ILKQAPPMNV	SSVCDNQPAV
	551				600
Penton5	TDHGTLPLRN	SIGGVQRVTI	TDARRRTCPY	VYKALGIVSP	RVLSSRTF*.
Penton2	TDHGTLPLRN	SIGGVQRVTI	TDARRRTCPY	VYKALGIVSP	RVLSSRTF*.
Penton3	TDHGTLPLRS	SIRGVQRVTV	TDARRRTCPY	VYKALGIVAP	RVLSSRTF*.
Penton12	TDHGTLPLRS	SISGVQRVTI	TDARRRTCPY	VYKALGIVSP	RVLSSRTF*.
Penton40	TDHGTLPLRS	SISGVQRVTI	TDARRRTCPY	VHKALGIVAP	KVLSSRTF*.
Penton17	TDHGTLPLRS	SIRGVQRVTV	TDARRRTCPY	VYKALGIVAP	RVLSSRTF*.
Pentonf10	VQQGVLPVKS	SLPGLQRVLI	TDDQRRPIPY	VYKSIATVQP	TVLSSATLQ*

Fiber17.Pep x Fiber2.Pep

		/ SEA	ID://
1	MSKRLRVEDDFNPVYPYGYARN.QNIPFLTPPFVSSDGFKNFPPGVLSLK	494 300	
	-:	50 ← SEQ!	D NO:12
-	LADPITIANGDVSLKVGGGLTLQE	73	
50	LADPITIANGDVSLKVGGGLTLQE		
51	LADPITIANGDVSLKVGGGTILQE ::: :: :: : : VSEPLDTSHGMLALKMGSGLTLDKAGNLTSQNVTTVTQPLKKTKSNISLD	100	
74	GSLTVDPKAPLOLANNKKLELVYVDPF	100	
	THE COLUMN TWO DESKI STATKER	150	
101	TSAPLTITSGALTVATTAPLIVTSGALSVQSQAFHIVQBBABBINA		
101	EVSANKLSLKVGHGLKILDDK	121	
	TSAPLTITSGALTVATTAPLIVTSGALSVQSQATTIVQSDASSTATIVGSDASST	200	
151	TVSDGKUADQTSAPESGSDSD1214111814		
122	SAGGLKDLIGKLVVLTGKGIGTE	144	
201	SAGGLKDLIGKLVVIIGKGIGIZ: : : : :. . : . :. GKIGIKISGPLQVAQNSDTLTVVIGPGVTVEQNSLRTKVAGAIGYDSSNN	250	
	•		
	•		
	NLONTD GSSRGIGISVRARE	164	
145	l:: :: :::		
301	:: . :: :: YNRGLYLFNASNNTKKLEVSIKKSSGLNFDNTAIAINAGKGLEFDTNTSE	350	
	CL TEDNICCYL VAWNPKYDTRT	185	
165	[1.][1.]:. []		
351	SPDINPIKTKIGSGIDYNENGAMITKLGAGLSFDNSGAITIGNKNDDKLT	400	
106	LWTTPDTSPNCRIDKEKDSKLTLVLTKCGSOILANVSLIVVSGKYQYIDH	235	
100	LWTTPDTSPNCRIDKEKDSKLTLVLTKCGSQULATV311	446	
401	LWTTPDPSPNCRIHSDNDCKFTLVLTKCGSQVLATVAALAVSGDLS	***	

FIGURE 7A

236	ATNPTLKSFKIKLLFDNKGVLLPSSNLDSTYWNFRSDNLTVSEAYKNAVE	285
447	:	496
286	FMPNLVAYPKPTTGSKKYAPDTIVONTSKI OOLA VODINITIOMENDEAD	222
497	: ::::: ::::: : :::: FMPNLLAYPKTQSQTAKNNIVSQVYLHGDKTKPMILTITLNGTSEST	543
334	SAYSITFEFVWNKE VARVEDETTERTERTER TO 266	
544		

0.511	1				50
8fiber	MTKRLRA	EDDFN	PVYPYGYARN	Q.NIPFLTPP	FVSSNGFQNF - SEA ID NO: 13
9fiber	MSKRLRV	EDDFN	PVYPYGYARN	Q.NIPFLTPP	FVSSDGFQNF - SEC ID NO: 14
15fiber	MSKRLRV	EDDFN	PVYPYGYARN	O.NIPFLTPP	FUSSINGRONE - SEA ID ANSIS
17fiber	MSKRLRV	EDDFN	PVYPYGYARN	O NTPFL/PDD	ENTERONOUS CALD AVAIL
2fiber	.MKRARP	SEDTFN	PVYPYDTETG	gan tagongg	FUSDNOROES
5f ibe r	TIMMARE	SEDIFN	PVYPYDIETG	ממית זיגוע טיויע ע	FUSDNICEOFS STAIR
4fiber	MSKSARG	WSDGFD	PVYPYDADND	RP CPSSTLP	SESSIVERORY SCA
40-1fiber	.MKRTRIE	DDFN	PVYPYD. 1199	מסגמדסעדמייי	EUCCIVIT OTAL
41fiber	.PIKKIKIE		PVYPYD. TPS	סס מוויסט ו פטיוי	EVECTOR OF
40-2fiber	. MARARCE	DDFN	PVIPIE HYN	- פסיידיאס וכווט	FASSNGI OFK
12fiber	.PICRORIUIA	EETERNIIIFN	PAABED BED	ססיתיאסטווציוי	ETICCNICI OFV
3fiber	MAKRARL	STSFN	PVYPYEDESS	SOH . PFINPG	FISPOGETQS SEA ID NO: 21
		•			Sea 10 NO: 12
	51		,		100
8fiber	PPGVLSLKLA	DPITIN.NQN	VSLKVGGGLT	LOEET	
9fiber	PPGVLSLKLA	DPIAIV.NGN	VSLKVGGGLT	LODGT	
15fiber	PPGVLSLKLA	DPIAIA.NGN	VSLKMGGGLT	LOEGT	
17fiber	PPGVLSLKLA	DPITIA.NGD	VSLKVGGGLT	LOE	
2fiber	PPGVLSLRVS	EPLDTS.HGM	LALKMGSGLT	LDKAGNLTSO	NVTTVTOPLK
5fiber	PPGVLSLRLS	EPLVTS.NGM	LALKMGNGLS	LDEAGNLTSO	NVTTVSPPLK
4fiber	PLGVLSLGPG	RPCHTK.NGE	ITLKLGEGVD	LDDSGKLTAN	TVNKATAPI.
40-1fiber	PPGVLALKYT	DPITTNAKHE	LTLKLGSNIT	LO. NGLLSA	
41fiber	PPGVLALKYT	DPITTNAKHE	LTLKLGSNIT	LE NGLISA	• • • • • • • • •
40-2fiber	PPGVLSLKYT	DPLTTK.NGA	LTLKLGTGLN	TDKNGDLSSD	ACVEVED PTT
12fiber	PPGVLALNYK	DPIVTE.NGT	LTLKLGDGTK	LNAOGOLTAS	MNTMMEDIA
3fiber	PNGVLSLKCV	NPLTTA.SGS	LOLKVGSGLT	VD.	1111111 1 2022 201
					* * * * * * * * * * * * * * * * * * * *
	101				150
8fiber					
9fiber					
15fiber					
17fiber					

FIGURE 8A

2fiber	KTKSNISLD	S LTITSGA	LTVATTAPLI	VTSGALS	OAPLTVODSK
5fiber	KTKSNINLEI	SAPLTVTSEA	LTVAAAAPLM	VAGNTLTMOS	OAPLTVHDSK
4fiber				SFFOOH	HFPL
40-1fiber	• • • • • • • • • •	• • • • • • • • • •			• • • • • • • • • •
41fiber	· · · · · · · · · · · · · · · · · · ·				• • • • • • • • • •
40-2fiber			LTLSYNAPFN		
12fiber 3fiber			LTLNTRAPLT		
2 Liber	• • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	
	151				200
8fiber					200
9fiber					
15fiber					
17fiber					
2fiber					LSI
5fiber					LST
4fiber					
40-1fiber					
41fiber					
40-2fiber		.NANNELSLL	IDAPLNADIG	TLRLRSDAPL	GLVDK.TLKV
12fiber	LGLATIAPLS	LDGGGNLGLN	LSAPLDVSNN	NLHLTTETPL	VVNSSGALSV
3fiber	• • • • • • • • • •	• • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •
	201				250
8fiber			GKLT	UNTEPPLH	250
9fiber			GKLT	VNADPPLO.	
15fiber			GNLT	VNTEPPLO	
17fiber			GSLT	VDPKAPLO	
2fiber	ATKGPITVSD	GKLALOTSAP	LSGSDSDTLT	VTASPPLTTA	TGSLGINMED
5 fib er	ATQGPLTVSE	GKLALQTSGP	LTTTDSSTLT	ITASPPLTTA	TGSLGIDLKE
4fiber		TWIP	LYTPKMENYP	YKFLPPLSIL	KSTI
40-1fiber		\dots TVPT \dots		VSPPLTNS	NNSLGLATSA
41fiber		\dots TVPT \dots		VSPPLTNS	NNSLGLATSA
40-2fiber	LFSSPLYLDN	NFLTLAIERP	LALSSNRAVA	LKYSPPLKIE	NENLTLSTGG
12fiber	ATADPISVRN	NALTLPTADP	LMVSSD.GLG	ISVTSPITVI	NGSLALSTTA
3fiber	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •			• • • • • • • • •
	251				200
8fiber		TALDADEDUT	D. NKLTLLA	CUCT CTT MY	300
9fiber	T.TINN KIG	TALDAFIDVI	D. NKLTLLA	CUCICIT TV	ETSTLIPGLVN
15fiber	I TWN RTG	TALDAPEDUI	GGKLTLLA	CHCICII TE	ETSIDEGEM
17fiber	LANNKKLE	LVYVDPFEVS	ANKLSLKV	CHCI.KII.DDK	CACCI-KDI-TC
2fiber	PIYVNNGKIG	IKISGPLOVA	QNSDTLTVVT	GPGVTVFONS	I.RTKVACATC
5fiber	PIYTONGKLG	LKYGAPLHVT	DDLNTLTVAT	GPGVTTNNTS	LOTKVTGALG
4fiber			LNTLVSAF	GSGLGLSGSA	LAVOLASPLT
40-1fiber	PIAVSANSLT	LATAAPLTVS	NNOLSINT	GRGLVITNNA	VAVNPTGALG
41fiber	PIAVSANSLT	LATAAPLTVS	NNOLSINA	GRGLVITNNA	LTVNPTGALG
40-2fiber	PFTVSGGNLN	LATSAPLSVO	NNSLSLGV	NPPFLITDSG	LAMDLGDGLA
12fiber	PLNSTGSTLS	LSVANPLTIS	Q. DTLTVST	GNGLOVSGSO	LVTRIGDGLT
3fiber	TTDGSLE	ENIKVNTPLT	KSNHSINLPI	GNGLQIEQNK	LCS
	301				354
8fiber					350
9fiber					
15fiber					
17fiber					
2fiber	YDSSNNMEIK	TGGGMRIN	NNLLILDVDY	PFDAQTKLRL	KLGOGPLYIN
				-	~

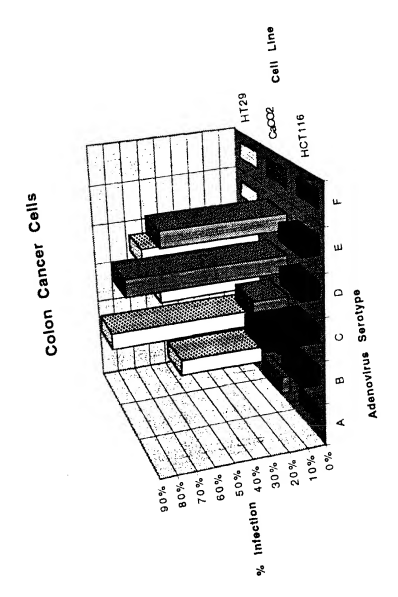
5fiber	FDSQGNMQLN	VA. JLRIDS	Q NRRLILDVS	PFDAQNQL	RLGOGPLFIN
4fiber 40-1fiber	FDDKG	*********	: : : : : : : : : : : : : : : : : : : :		• • • • • • • • • • • • • • • • • • • •
41fiber	TANTICAL OLM	AAGGMRVDG	A N. LILHVA	PFEAINQLTL	R
40-2fiber	TOO CRITIN	AAGGMRVDG	A N. LILHVAY	PFEAINQLTL	R
12fiber	TOG - DUTITIN	LGPGLQMSN	G AITL	ALDAALPL	<u></u> Q
3fiber	PDN.GVERVN	VAGGMRTSG	G RIILDVNY	PFDASNNLSL	RRGLGLIYNQ
TIMET		• • • • • • • • •		• • • • • • • • • • •	
	351				400
8fiber					400
9fiber				• • • • • • • • • • •	TEV VETGRET
15fiber				••••••	TDAADIGVGT
17fiber				• • • • • • • • • •	TTA ATTICKOT
2fiber	ASHNUDINYN	RGI.YT.FNAS	N NTKKLEVSIK	WCCCI MEDAM	VENATIONET
5fiber	SAHNI DINYN	KCI.VI.FTAC	N NSKKLEVNLS	. WOOGTMEDMI.	ALAINAGKGL
4fiber	NTKTTLN	PCT.Hymrych	AIESNIS	TWENTER DW.I.	ALAINAGIGL
40-1fiber		remin 116D	HIESMIS	MAYCTVERIG	ALATNIGKGS
41fiber		• • • • • • • • •	· · · · · · · · · · · · · · · · · · ·	• • • • • • • • • • •	• • • • • • • • •
40-2fiber	VKNN		·	• • • • • • • • • • • •	01.01.5
12fiber	CUMMI			• • • • • • • • • • •	QLQLRIGS
3fiber	D144	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •	NLTTDIST
TIMEL	• • • • • • • • • •	• • • • • • • • • •		• • • • • • • • • • •	• • • • • • • • • •
	401				450
8fiber		NICVRV	3 E	מממז פ	שונג זרוטאכוטא
9fiber	GTESTDNGG	TVCVRV	G E	CCCTC	ENDIDGE TANE
15fiber	GTDTTDNGG.	STRVRV	E	CCCI C	EMEY COLLINE LIMINGODO VAL
17fiber	GTENLONTDG	SSRGTGTSVI	A	בענטטט	FINEAGDLIVAE
2fiber	EFDINTSESP	DINPIKTKI	SGIDYNENGA	MTTTKI CACI C	FUNDGITAW
5fiber	EFG. SPNAP	NTNPLKTKT(HGLEFDSNKA	MIDRI CACI C	FDCDCATTIG
4fiber	REGUSSTETC	UNINIAVDIOU	· · · · · · · · · · · · · · · · · · · ·	MALCOCT C	FDSTGATTVG
40-1fiber	in Gibbillio	TI	NGLEVINGGK	THE CCOLO	FDSTGAIMAG
41fiber	• • • • • • • • • •		NOTE MICCON	TWANTYZGTŐ	FUNNGRITIS
40-2fiber	ACAT TMCCTM	OUL STATES	NGLEVTSGGK	LINVKLGSGLQ	FDSNGRIAIS
12fiber	ERGI MEGGA PPRTTTPGALI	OT STANISMENTS	KGLAIENNS.	LVVKLGNGLR	FDSWGSIAVS
3fiber	ENGINE SGN.	QIALMAC	QGLTFNNGQ.	LKVKLGAGLI	FDSNNNIALG
PITDEL		• • • • • • • • • • • • • • • • • • • •		KLGNGLT	FDSSNSIALK
	451				500
8fiber		.RTLWTTPDT	SPNCRID	ODKDSKLSLV	L/PKCGGATLA
9fiber	NKKEDK	RTLWTTPD	SPNCKID	ODKDSKIMIA	TWCGSOTTW
15fiber	NKKEDM	RTLWTTPDI	SPNCKII	EDKDGKI MI 1	TURCOSÕTTA
17fiber	NPKYDT	ירופידיניין זיין	SPNCRID	KEKDGKI WI 11	T WACGEOTT &
2fiber	NKNDDK	זרוטיזידיש, זיף, ז	SPNCRIH	CDMIA:ABUL 43	T WACCCOAL *
5fiber	NKNNDK	ז מסידים שינים ב	SPNCRIN	PDMICKLITA	T WYCGSQVLA
4fiber	NKDADA	THATTAINTEE.	CDNCCT!	AENDAKETEV	LTKCGSQ1LA
40-1fiber	MDIOMOGRAMO	THATPATTADE	SPNCQIL	AENDAKLTLC	LIMCUSQILA
41fiber	MCMBMbware	TUMPTOTO .	TPNCSIY	ETODANLFLC	LTKNGAHVLG
40-2fiber	MOMENTA	TITIMSIS. I	TPNCSIY	ETQUANLFLC	LTKNGAHVLG
12fiber	PITITP.	TILWITADI	SPNATFY	ESLDAKVWLV	LVKCNGMVNG
	SSSNTPYDP.	LTLWTTPDE	PPNCSLI	QELDAKLTLC	LTKNGSIVNG
3fiber	NN	TLWTGPK	' EANCIIEYGK	QNPDSKLTLI	LVKNGGIVNG
	501				
8fiber		VVT TRIKINGS	X 7 7/0		550
9fiber	MUSILIVVAGR	IVT TWWWIW!	ALKGFTIK	LLFDKNGVLM	ESSN
	NVSLIVVDGK	TYTTUNNIO	·ALKGFTIK	LLFDENGVLM	ESSN
15fiber	SVSLLVVKGK	FSNINNTTNE	NEADKQITVK	LLFDANGVLK	QGST
17fiber	NVSLIVVSGK	YQYIDHATNE	TLKSFKIK	LLFDNKGVLL	PSSN.
2fiber	TVAALAV.S.	GDLSSM	TGTVASVSIF	LRFDONGVLM	ENCC
5fiber	TVSVLAV.K.	GSLAPI	SGTVQSAHLI	IRFDENGVLL	NNSF
			•		

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4fiber TVSVLVVRS. .. GNLNPI TGTVSSAQVF LRFDANGV
40-1fiber TITIKGLKGA LREMNDNA.....LSVK LPFDNQGNLL NCA.....
   41fiber TITIKGLKGA LREMHDNA. LSLK LPFDNQGNLL NCA.

-2fiber TISIKAQKGT LL. KPTASF ... ISFV MYFYSDGTWR KNYPVFDNEG
12fiber IVSLVGVKGN LLNIQSTTTT ... VGVH LVFDEQGRLI TSTP....T
40-2fiber
     3fiber YVTLMGASDY VNTLFKNKNV .....SINVE LYFDATGHIL PDSSSLKTDL
    8fiber ..LGKSYWNF RNQNSIMSTA YEKAIGFMPN LVAYPKPTTG SKKY...ARD
    9fiber ..LGKSYWNF RNENSIMSTA YEKAIGFMPN LVAYPKPTAG SKKY...ARD
   15fiber ..MDSSYWNY RSDNSNLSQP YKKAVGFMPS KTAYPKQTKP TNKEISQAKN
17fiber ..LDSTYWNF RSDNLTVSEA YKNAVEFMPN LVAYPKPTTG SKKY..ARD
2fiber .LKKHYWNF RNGNSTNANP YTNAVGFMPN LLAYPKTQSQ T....AKN
    5fiber ...DPEYWNF RNGDLTEGTA YTNAVGFMPN LSAYPKSHGK T.....AKS
4fiber .TSKKYWGY KQGDSIDGTP YTNAVGFMPN STAYPKTQSS T...TKN
40-1fiber .LESSTWRY QETNAVA...SNALTFMPN STVYPRNKTA D...PGN
41fiber .LESSTWRY QETNAVA...SNALTFMPN STVYPRNKTA H...PGN
40-2fiber ILANSATWGY RQGQSANTN. VSNAVEFMPS SKRYPNEKGS E...VQN
   12fiber ALVPQASWGY RQGQSVSTNT VTNGLGFMPN VSAYPRPNAS E...AKS
3fiber ELKYKQTADF .....SARGFMPS TTAYPFVLPN AGTH..NEN
    8fiber IVYGNIYLGG KPHQ..PVTI KTTFNQETG. ....CEYS ITFDFSWAKT 9fiber IVYGNIYLGG KPDQ..PVTI KTTFNQETG. .....CEYS ITFDFSWAKT
   15fiber KIVSNVYLGG KIDQ..PCVI IISFNEEAD. .....SDYS IVFYFKWYKT
   17fiber IVYGNIYLGG LAYQ. PVVI KVTFNEEAD. .....SAYS ITFEFVWNKE
    2fiber NIVSQVYLHG DKTK..PMIL TITLNGTSES TETSEVSTYS MSFTWSWESG
5fiber NIVSQVYLNG DKTK..PVTL TITLNGTQET GDTT.PSAYS MSFSWDWSGH
    4fiber NIVGQVYMNG DVSK..PMLL TITLNGTDDT T....SAYS MSFSYTWTNG
40-1fiber MLI...... QISP..NITF SVVYNEINS......GYA FTFKW.SAEP
41fiber MLI...QISP..NITF SVVYNEINS.....GYA FTFKW.SAEP
40-2fiber MALTYTFLQG DPNM..AISF QSIYN..HA...IEGYS LKFTW.RVRN
   12fiber OMVSLTYLOG DTSK..PITM KVAFNGITS.....LNGYS LTFMW.SGLS
    3fiber YIFGOCYYKA SDGALFPLEV TVMLNKRLPD SRTSYVMTFL WSLNAGLAPE
    8fiber .YVNVEFETT SFTFSYIAOE *
    9fiber .YVNVEFETT SFTFSYIAQE *
  15fiber .YENVQFDSS SFNFSYIAQE * 17fiber .YARVEFETT SFTFSYIAQQ *
    2fiber KYTTETFATN SYTFSYIAQE ...
5fiber NYINEIFATS SYTFSYIAQE *
4fiber SYIGATFGAN SYTFSYIAQQ *
40-1fiber ...GKPFHPP TAVFCYITEQ *
41fiber ...GKPFHPP TAVFCYITEQ *
40-2fiber ...NERFDIP CCSFSYVTEQ
  12fiber NYINOPFSTP SCSFSYITOE *.
3fiber T.TQATLITS PFTFSYIRED D*
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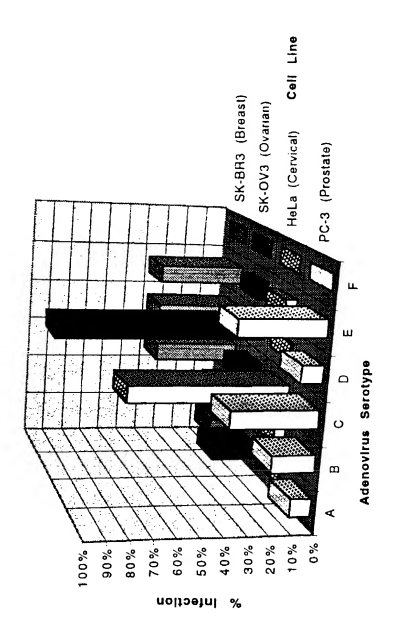
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Fleure 4

Cancer Cell Lines



LXAMINE 10

Interna al Application No PCT/US 97/21494

a. classif IPC 6	FICATION OF SUBJECT MATTER C12N15/86 A61K48/00								
According to	o International Patent Classification (IPC) or to both national classific	eation and IPC							
B. FIELDS	SEARCHED								
IPC 6	cumentation searched (classification system followed by classificat C12N A61K C07K								
Documentat	tion searched other than minimum documentation to the extent that	such documents are included in the fields sea	rched						
Electronia d	ata base consulted during the international search (name of data b	ase and, where practical, search terms used)							
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT								
Category *	Citation of document, with indication, where appropriate, of the re	elevant passages	Relevant to claim No.						
A	P.W. ROELVINK ET AL.: "Compara- analysis of adenovirus fiber-ce interaction: Ad2 and Ad9 utilize cellular fiber receptor but use	ll e the same different	1-13						
	binding strategies for attachment" JOURNAL OF VIROLOGY, vol. 70, no. 11, November 1996, AMERICAN SOCIETY FOR MICROBIOLOGY US, pages 7614-7621, XP002062100 see page 7620, last paragraph								
A	WO 96 26281 A (GENVEC INC ;CORN FOUNDATION INC (US)) 29 August see example 7	ELL RES 1996 -/	1,4,6-8, 10,11						
X Fur	rther documents are listed in the continuation of box C.	X Patent family members are listed	in annex.						
"A" doour consi "E" earlier filing "L" doour whiol citati "O" doour other	categories of cited documents : ment defining the general state of the art which is not idered to be of particular relevance r document but published on or after the international	"I" later document published after the interpriority date and not in conflict with cited to understand the principle or the invention of the principle of the invention of the cannot be considered novel or cannot involve an inventive step when the deciment of particular relevance; the cannot be considered to involve an indocument is combined with one or ments, such combined with one or ments, such combination being obviding the art. "&" document member of the same patent.	in the appearant of the cory underlying the cory underlying the considered to cournent is taken alone claimed invention the core other such doout-out to a person skilled						
Date of the	e actual completion of the international search 14 April 1998	Date of mailing of the international se							
	d mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Cupido, M							

Interr nal Application No PCT/US 97/21494

		PCT/US 9//21494
<u> </u>	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	Relevant to claim No.
Category °	Citation of document, with indication, where appropriate, of the relevant passages	TORY SUIT OF SIGNIFIES.
A	J. GALL ET AL: "Adenovirus type 5 and 7 capsid chimera: Fiber replacement alters receptor tropism without affecting primary immune neutralization epitopes" JOURNAL OF VIROLOGY., vol. 70, no. 4, April 1996, pages 2116-2123, XP002050655 see the whole document	1,4,6-8, 10,11
P,X	see the whole document WO 97 12986 A (CORNELL RES FOUNDATION INC) 10 April 1997 see page 15, line 1 - line 7	1,2,13

Ir. ational application No.

PCT/US 97/21494

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of Iirst sheet)
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2. Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is reatricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

information on patent family members

Interr nat Application No PCT/US 97/21494

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 9626281 A	29-08-96	AU 4980496 A CA 2213343 A EP 0811069 A	11-09-96 29-08-96 10-12-97
WO 9712986 A	10-04-97	NONE	

F1G. 8C-1

RLGQGPLFIN R	400 % TLVVLTGKGI TLVVLTGKGI TLVVLTGKGL KLVVLTGKGL AIAINAGDGL AIAINAGDGL AIAINAGDGL AIATNIGKGSQLQLRIGSNLTTDIST
PFDAQNQLNL RLGQGPLFIN PFEAINQLTL R PFEAINQLTL R ALDAALPL OPFDASNNLSL RRGLGLIYNQ	KSSGLNFDNT TAKGLMFDAT WAKGIKFEDG
NRRLILDVSY N. LILHVAY N. LILHVAY A. ITL R. IILDVNY	NTKKLEVSIK NSKKLEVNLS . IESNIS
VAGGLRIDSQ AAGGMRVDGA AAGGMRVDGA LGPGLQMSNG VAGGMRTSGG	RGLYLFNASN KGLYLFTASN RGLHVTTGDA
FDSQGNMQLN FDDKG FNNTGALQLN FNNTGALQLN LGG.SKLIIN FDN.GVMKVN	351 ASHNLDINYN SAHNLDINYNNIKITLN YKNN
5fiber 4fiber 40-1fiber 41fiber 40-2fiber 12fiber	8 fiber 9 fiber 15 fiber 17 fiber 2 fiber 4 fiber 40-1 fiber 40-2 fiber 12 fiber 3 fiber 3 fiber

F16.8C-2

450	GGGLS FNDNGDLVAF	GGGLS FNNDGDLVAF	GGGLS FNEAGDLVAF	REGLT FDNDGYLVAW	FDNSGAITIG	FDSTGAITVG	KLGSGLS FDSTGAIMAG	LNVKLGSGLQ FDNNGRITIS	LNVKLGSGLQ FDSNGRIAIS	LVVKLGNGLR FDSWGSIAVS	LRVKLGAGLI FDSNNNIALG	KLGNGLT FDSSNSIALK
	···· GGGLS	···· GGGLS	···· GGGLS	REGLT	MITKLGAGLS FDNSGAITIG	MVPKLGTGLS FDSTGAITVG				LVVKLGNGLR	LRVKLGAGLI	KLGNGLT
	NICVRVG E	田	田		SGIDYNENGA	NTNPLKTKIG HGLEFDSNKA	•	LE NGLEVINGGK	NGLEVTSGGK	KGLAIENNS.	QGLTFNNGQ.	•
	NICVRVG	TVCVRVG	SIRVRVG	SSRGIGISVR	DINFIKTKIG	NTNPLKTKIG	VNNAYPIQV.	IE	E	OTLNVNANTS KGLAIENNS.	QIALNAG QGLTFNNGQ.	
401	GTDLSNNGG.	GTESTDNGG.	GTDTTDNGG.	GTENLQNTDG	EFDTNTSESP	EFGSPNAP	RFGTSSTETG	•		ASALIMSGVT	EKGLMFSGN.	•
	8fiber	9fiber	15fiber	17fiber	2fiber	5fiber	4fiber	40-1fiber	41fiber	40-2fiber	12fiber	3fiber

23/28

SUBSTITUTE SHEET (RULE 26)

F1G. 8C-3

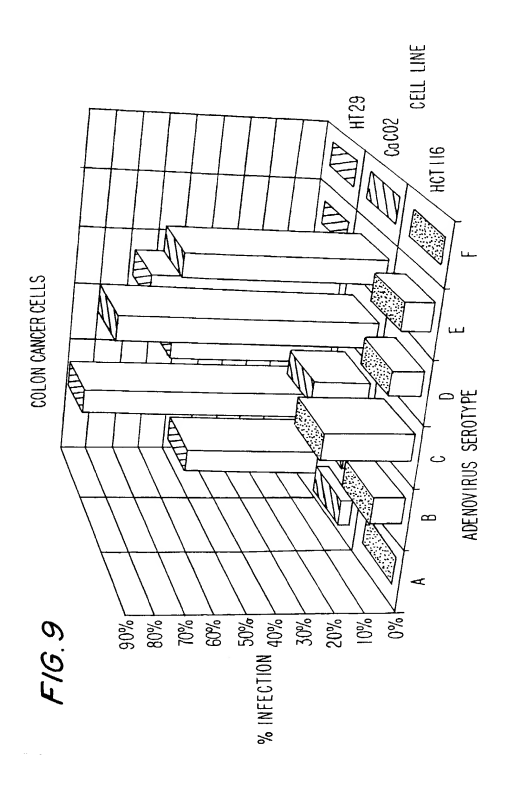
SPNCRID QDKDSKLSLV SPNCKID QDKDSKLTLV SPNCKII EDKDSKLTLI	SPNCRID KEKDSKLTLV SPNCRIH SDNDCKFTLV SPNCRLN AEKDAKLTLV SPNCQIL AENDAKLTLC	TPNCSIY TPNCSIY	PPNCSLI QELDAKLTLC EANCIIEYGK QNPDSKLTLI	ALKGFTIK LLFDKNGVLM ESSN	NEADNOIL ON DELEGNICOLES. TLKSFKIK LLFDNKGVLL TGTVASVSIF LRFDQNGVLM SGTVQSAHLI IRFDENGVLL
RTLWTTPDT. RTLWTTPDT. RTLWTTPDP	. RTLWTTPDT . LTLWTTPDP . LTLWTTPAP . LTLWTTPDP	LTTIWSIS.P LTTIWSIS.P .TTLWTTADP	.LTLWTTPDP .TLWTGPKP	YKIINNNTNP YKIINNNTOP	YQYIDHATNP YQYIDHATNP GDLSSM
451 NKKEDK NKKEDK			SSSNTPYDP.	501 NVSLIVVAGR NVSLIVVDGK	
8fiber 9fiber 15fiber	17fiber 2fiber 5fiber 4fiber	40-1fiber 41fiber 40-2fiber	12fiber 3fiber	8fiber 9fiber	15fiber 17fiber 2fiber 5fiber

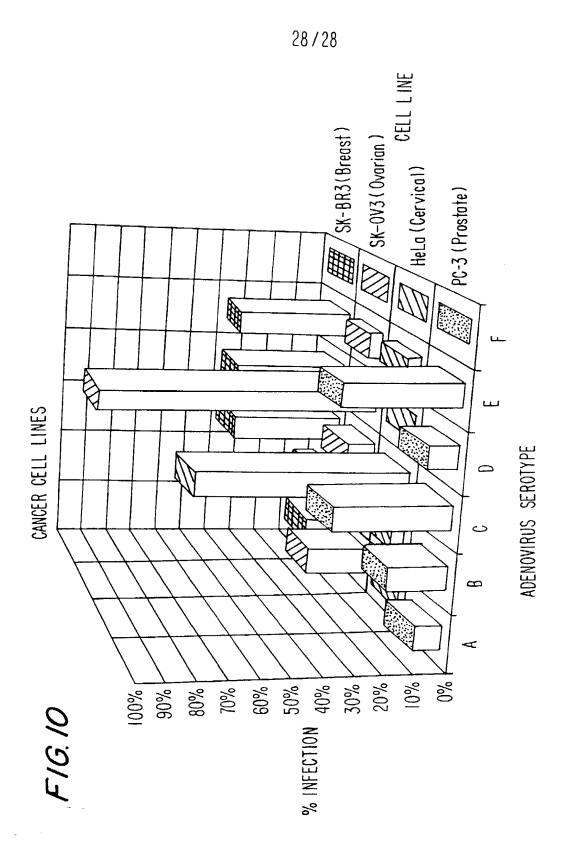
F1G. 8D-1

TEHS NCA NCA KNYPVFDNEG TSTPT PDSSSLKTDL	600 SKKYARD SKKYARD TNKEISQAKN	SKKYARD TAKS	TTKN DPGN HPGN	EVQN EAKS AGTHNEN
LRFDANGVLL LPFDNQGNLL LPFDNQGNLL MYFYSDGTWR LVFDEQGRLI LVFDEQGRLI	LVAYPKPTTG LVAYPKPTAG KTAYPKQTKP	LVAYPKPTTG LLAYPKTQSQ LSAYPKSHGK	STAYPRIVSS STVYPRNKTA STVYPRNKTA	SKRYPNEKGS VSAYPRPNAS TTAYPFVLPN
TGTVSSAQVFLSVKLSLKISFVSINVE	YEKAIGFMPN YEKAIGFMPN YKKAVGFMPS	YKNAVEFMPN YTNAVGFMPN YTNAVGFMPN	YTNAVGFMPN SNALTFMPN SNALTFMPN	VSNAVEFMPS VTNGLGFMPN SARGFMPS
LREMNDNA LREMHDNA LLKPTASF LLNIQSTTTT	RNQNSIMSTA RNENSIMSTA RSDNSNLSQP	RSDNLTVSEA RNGNSTNANP RNGDLTEGTA	KQGDSIDGTP QETNAVA QETNAVA	RQGQSANTN. RQGQSVSTNT
TVSVLVVRS. TITIKGLKGA TITIKGLKGA TISIKAQKGT IVSLVGVKGN	551 LGKSYWNF LGKSYWNF	LDSTYWNF LKKHYWNF LDPEYWNF	. TSKKYWGY . LESSTWRY . LESSTWRY	ILANSATWGY ALVPQASWGY ELKYKQTADF
4fiber 40-1fiber 41fiber 40-2fiber 12fiber 3fiber	8fiber 9fiber 15fiber	17fiber 2fiber 5fiber	4tiber 40-1fiber 41fiber	40-2fiber 12fiber 3fiber

25/28

650	CEYS ITFDFSWAKT		SAYS ITFEFVWNKE		•	_	GYA FTFKW.SAEP	GYA FTFKW.SAEP	IEGYS LKFTW.RVRN	LNGYS LTFMW.SGLS	SRTSYVMTFL WSLNAGLAPE							110.0UZ							
	KTTFNQETG. KTTFNQETG.	IISFNĒEAD.	KVTFNEEAD.	TITLNGTSES	TITLNGTQET	TITLNGTDDT	SVVYNEINS.	SVVYNEINS.	QSIYNHA.	KVAFNGITS.	TVMLNKRLPD	0	7/9	*		*	*	•	*		*		*	*	μΩ*
	KPHQPVTI KPDOPVTI	KIDQPCVI	LAYQPVVI	DKTKPMIL	DKTKPVTL	DVSKPMLL	QISPNITF	QISPNITF	DPNMAISF	DTSKPITM	SDGALFPLEV		ט	SFTFSYIAQE	SFTFSYIAQE	SFNFSYIAQE	SFTFSYIAQQ	SYTESYIAQE			TAVFCYITEQ	TAVFCYITEQ	CCSFSYVTEQ	SCSFSYITQE	PFTFSYIRED
601	IVYGNIYLGG	KIVSNVYLGG	IVYGNIYLGG	NIVSQVYLHG	NIVSQVYLNG	NIVGQVYMNG	MLI	MLI	MALTYTFLQG	OMVSLTYLQG	YIFGQCYYKA	,	65.I	YVNVEFETT.	YVNVEFETT	YENVQFDSS	YARVEFETT	KYTTETFATN	NYINEIFATS	SYIGATFGAN	GKPFHPP	GKPFHPP	NERFDIP	NYINOPESTP	T. TQATLITS
	8fiber 9fiber	15fiber		iber	iber	iber		41fiber	40-2fiber	12fiber	3fiber			8fiber	9fiber	15fiber	17fiber	2fiber	5fiber	4fiber	_	$\overline{}$	40-2fiber		3fiber





Interna al Application No PCT/US 97/21494

	•		700 31722151
A CLASSII IPC 6	FICATION OF SUBJECT MATTER C12N15/86 A61K48/00		
According to	o International Patent Classification (IPC) or to both national class	ification and IPC	
B. FIELDS	SEARCHED		
Minimum do IPC 6	ocumentation searched (classification system followed by classifi C12N A61K C07K	oation symbols)	
Documenta	tion searched other than minimum documentation to the extent th	at such documents are included in (he fields searched
Electronic d	lata base consulted during the international search (name of date	base and, where practical, search	terms used)
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
A	P.W. ROELVINK ET AL.: "Comparanalysis of adenovirus fiber-cinteraction: Ad2 and Ad9 utilicellular fiber receptor but us binding strategies for attachm JOURNAL OF VIROLOGY, vol. 70, no. 11, November 1996 SOCIETY FOR MICROBIOLOGY US, pages 7614-7621, XP002062100 see page 7620, last paragraph	ell ze the same e different ent ^m	1-13
A	WO 96 26281 A (GENVEC INC ;COR FOUNDATION INC (US)) 29 August see example 7	NELL RES 1996 -/	1,4,6-8, 10,11
X Fur	ther documents are listed in the continuation of box C.	X Patent family membe	rs are listed in annex.
"A" doourn "E" earlier filing "L" docum which obtails "O" docum other "P" docum later	ategories of cited documents: ment defining the general state of the art which is not detect to be of particular relevance of coursent but published on or after the international date each which may throw doubts on priority claim(s) or his cited to establish the publication date of another on or or ther special reason (as specified) ment referring to an oral disclosure, use, exhibition or means enter published prior to the international filing date but than the priority date claimed sectual completion of the international search	or priority date and not in cited to understand the privention "X" document of particular relocannot be considered no involve an inventive step "Y" document of particular relocannot be considered to document is combined with the control of the combination in the art. "&" document member of the	
	14 April 1998	123.04.98	
Name and	mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijawijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Cupido, M	

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Interr nal Application No PCT/US 97/21494

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	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	Relevant to plaim No.
ategory °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to olaim No.
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,,X	WO 97 12986 A (CORNELL RES FOUNDATION INC) 10 April 1997 see page 15, line 1 - line 7	1,2,13
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Ir. ,ational application No.

PCT/US 97/21494

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Remark on Protest The additional search fees were accompanied by the applicant's prot No protest accompanied the payment of additional search fees.	est.

information on patent family members

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Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 9626281 A	29-08-96	AU 4980496 A CA 2213343 A EP 0811069 A	11-09-96 29-08-96 10-12-97
WO 9712986 A	10-04-97	NONE	